

09/700259

532 Rec'd PCT/PTO 14 NOV 2000

IN THE UNITED STATES PATENT AND TRADEMARK OFFICE
REQUEST FOR FILING NATIONAL PHASE OF
PCT APPLICATION UNDER 35 U.S.C. 371 AND 37 CFR 1.494 OR 1.495

To: Hon. Commissioner of Patents
 Washington, D.C. 20231

TRANSMITTAL LETTER TO THE UNITED STATES
 DESIGNATED/ELECTED OFFICE (DO/EO/US)

Atty Dkt: PM 275388 /P-75076US
M# /Client Ref.

From: Pillsbury Madison & Sutro LLP, IP Group: Date: November 14, 2000

This is a **REQUEST** for **FILING** a PCT/USA National Phase Application based on:

1. International Application	2. International Filing Date	3. Earliest Priority Date Claimed
<u>PCT/GB99/01362</u> <u>↑ country code</u>	17 MAY 1999 Day MONTH Year	15 MAY 1998 Day MONTH Year (use item 2 if no earlier priority)

4. Measured from the earliest priority date in item 3, this PCT/USA National Phase Application Request is being filed within:

(a) 20 months from above item 3 date (b) 30 months from above item 3 date,

(c) Therefore, the due date (unextendable) is November 15, 2000

5. Title of Invention GENE THERAPY VECTORS AND THEIR USE IN ANTITUMOUR THERAPY

6. Inventor(s) MARGISON, Geoffrey P., et al

7. Applicant herewith submits the following under 35 U.S.C. 371 to effect filing:

7. Please immediately start national examination procedures (35 U.S.C. 371 (f)).

8. A copy of the International Application as filed (35 U.S.C. 371(c)(2)) is transmitted herewith (file if in English but, if in foreign language, file only if not transmitted to PTO by the International Bureau) including:

- Request;
- Abstract;
- ____ pgs. Spec. and Claims;
- ____ sheet(s) Drawing which are informal formal of size A4 11"

9. A copy of the International Application has been transmitted by the International Bureau.

10. A translation of the International Application into English (35 U.S.C. 371(c)(2))

- is transmitted herewith including: (1) Request; (2) Abstract;
 (3) ____ pgs. Spec. and Claims;
 (4) ____ sheet(s) Drawing which are:
 informal formal of size A4 11"
- is not required, as the application was filed in English.
- is not herewith, but will be filed when required by the forthcoming PTO Missing Requirements Notice per Rule 494(c) if box 4(a) is X'd or Rule 495(c) if box 4(b) is X'd.
- Translation verification attached (not required now).

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RE: USA National Filing of PCT /GB99/01362

11. **PLEASE AMEND** the specification before its first line by inserting as a separate paragraph:
 a. --This application is the national phase of international application PCT/GB99/01362 filed May 17, 1999 which designated the U.S.--
 b. --This application also claims the benefit of U.S. Provisional Application No. 60/ , filed .

12. Amendments to the claims of the International Application **under PCT Article 19 (35 U.S.C. 371(c)(3))**, i.e., **before 18th month from first priority date above in item 3, are transmitted herewith (file only if in English) including:**

13. PCT Article 19 claim amendments (if any) have been transmitted by the International Bureau

14. Translation of the amendments to the claims **under PCT Article 19 (35 U.S.C. 371(c)(3))**, i.e., of **claim amendments made before 18th month, is attached (required by 20th month from the date in item 3 if box 4(a) above is X'd, or 30th month if box 4(b) is X'd, or else amendments will be considered canceled)**.

15. **A declaration of the inventor (35 U.S.C. 371(c)(4))**
 a. is submitted herewith Original Facsimile/Copy
 b. is not herewith, but will be filed when required by the forthcoming PTO Missing Requirements Notice per Rule 494(c) if box 4(a) is X'd or Rule 495(c) if box 4(b) is X'd.

16. **An International Search Report (ISR):**
 a. Was prepared by European Patent Office Japanese Patent Office Other
 b. has been transmitted by the international Bureau to PTO.
 c. copy herewith (pg(s).) plus Annex of family members (pg(s).).

17. **International Preliminary Examination Report (IPER):**
 a. has been transmitted (if this letter is filed after 28 months from date in item 3) in English by the International Bureau with Annexes (if any) in original language.
 b. copy herewith in English.
 c. 1 IPER Annex(es) in original language ("Annexes" are amendments made to claims/spec/drawings during Examination) including attached amended:
 c. 2 Specification/claim pages # claims #
 Dwg Sheets #
 d. Translation of Annex(es) to IPER **(required by 30th month due date, or else annexed amendments will be considered canceled)**.

18. **Information Disclosure Statement including:**
 a. Attached Form PTO-1449 listing documents
 b. Attached copies of documents listed on Form PTO-1449
 c. A concise explanation of relevance of ISR references is given in the ISR.

19. **Assignment document and Cover Sheet for recording are attached. Please mail the recorded assignment document back to the person whose signature, name and address appear at the end of this letter.**

20. Copy of Power to IA agent.

21. **Drawings (complete only if 8d or 10a(4) not completed):** sheet(s) per set: 1 set informal; Formal of size A4 11"

22. Small Entity Status is **Not claimed** is claimed **(pre-filing confirmation required)**
 22(a) (No.) Small Entity Statement(s) enclosed (since 9/8/00 Small Entity Statements(s) not essential to make claim)

23. **Priority** is hereby claimed under 35 U.S.C. 119/365 based on the priority claim and the certified copy, both filed in the International Application during the international stage based on the filing in (country) Great Britain of:

	Application No.	Filing Date	Application No.	Filing Date
(1)	9810423.5	15 MAY 1998	(2)	
(3)			(4)	
(5)			(6)	

a. See Form PCT/IB/304 sent to US/DO with copy of priority documents. If copy has not been received, please proceed promptly to obtain same from the IB.
 b. Copy of Form PCT/IB/304 attached.

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RE: USA National Filing of PCT/GB99/01362

24. Attached:

25. Preliminary Amendment:

25.5 Per Item 17.c2, cancel original pages #_____, claims #_____, Drawing Sheets #26. Calculation of the U.S. National Fee (35 U.S.C. 371 (c)(1)) and other fees is as follows:Based on amended claim(s) per above item(s) 12, 14, 17, 25, 25.5 (hilite)

Total Effective Claims	minus 20 =	x \$18/\$9 =	\$0	966/967
Independent Claims	minus 3 =	x \$80/\$40 =	\$0	964/965
If any proper (ignore improper) Multiple Dependent claim is present,		add\$270/\$135	+0	968/969

BASIC NATIONAL FEE (37 CFR 1.492(a)(1)-(4)): →→ BASIC FEE REQUIRED, NOW →→→A. If country code letters in item 1 are not "US", "BR", "BB", "TT", "MX", "IL", "NZ", "IN" or "ZA"

See item 16 re:

1. Search Report was <u>not prepared</u> by EPO or JPO -----	add\$1000/\$500	960/961
2. Search Report was prepared by EPO or JPO -----	add\$860/\$430	+430

SKIP B, C, D AND E UNLESS country code letters in item 1 are "US", "BR", "BB", "TT", "MX", "IL", "NZ", "IN" or "ZA"

→ <input type="checkbox"/> B. If <u>USPTO</u> did not issue <u>both</u> International Search Report (ISR) and (if box 4(b) above is X'd) the International Examination Report (IPER), -----	add\$970/\$485	+0	960/961
→ <input type="checkbox"/> C. If <u>USPTO</u> issued ISR but not IPER (or box 4(a) above is X'd), -----	add\$710/\$355	+0	958/959
→ <input type="checkbox"/> D. If <u>USPTO</u> issued IPER but IPER Sec. V boxes <u>not all</u> 3 YES, -----	add\$690/\$345	+0	956/957
→ <input type="checkbox"/> E. If international preliminary examination fee was paid to <u>USPTO</u> and Rules 492(a)(4) and 496(b) <u>satisfied</u> (IPER Sec. V <u>all</u> 3 boxes YES for <u>all</u> claims), -----	add \$100/\$50	+0	962/963

27.	SUBTOTAL =	\$430
28. If Assignment box 19 above is X'd, add Assignment Recording fee of ---\$40	+0	(581)
29. Attached is a check to cover the -----	TOTAL FEES	\$430

Our Deposit Account No. 03-3975

Our Order No. 40432

275388

C#

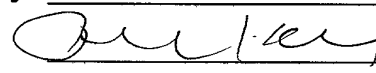
M#

CHARGE STATEMENT: The Commissioner is hereby authorized to charge any fee specifically authorized hereafter, or any missing or insufficient fee(s) filed, or asserted to be filed, or which should have been filed herewith or concerning any paper filed hereafter, and which may be required under Rules 16-18 and 492 (missing or insufficient fee only) now or hereafter relative to this application and the resulting Official document under Rule 20, or credit any overpayment, to our Account/Order Nos. shown above for which purpose a duplicate copy of this sheet is attached.

This CHARGE STATEMENT does not authorize charge of the issue fee until/unless an issue fee transmittal form is filed

Pillsbury Madison & Sutro LLP
Intellectual Property Group

1100 New York Avenue, NW
Ninth Floor
Washington, DC 20005-3918
Tel: (202) 861-3000
Atty/Sec: PNK/sdm

By Atty: <u>Paul N. Kokulis</u>	Reg. No. <u>16773</u>
Sig: 	Fax: (202) 822-0944
	Tel: (202) 861-3503

NOTE: File in duplicate with 2 postcard receipts (PAT-103) & attachments.

Applicant/Patentee: GEOFFREY PAUL MARGISON, ET AL
Serial No./Patent No.: 09/700,259
Filed or Issued: November 14, 2000
For: GENE THERAPY VECTORS AND THEIR USE IN ANTITUMOUR THERAPY

VERIFIED STATEMENT (DECLARATION) CLAIMING SMALL ENTITY STATUS (37 CFR 1.9(f) and 1.27(b)) - INDEPENDENT INVENTOR

As a below named inventor, I hereby declare that I qualify as an independent inventor as defined in 37 CFR 1.9(c) for purposes of paying reduced fees under Section 41(a) and (b) of Title 35, United States Code, to the Patent and Trademark Office with regard to the invention entitled GENE THERAPY VECTORS AND THEIR USE IN ANTITUMOUR THERAPY described in.

[] the specification filed herewith
[X] application serial no. 09/700,259,
 filed November 14, 2000
[] patent no. _____, issued _____

I have not assigned, granted, conveyed or licensed and am under no obligation under contract or law to assign, grant, convey or license, any rights in the invention to any person who could not be classified as an independent inventor under 37 CFR 1.9(c) if that person had made the invention, or to any concern which would not qualify as a small business concern under 37 CFR 1.9(d) or a non-profit organization under 37 CFR 1.9(e).

Each person, concern or organization to which I have assigned, granted, conveyed, or licensed or am under an obligation under contract or law to assign, grant, convey, or license any rights in the invention is listed below:

[] no such person, concern, or organization
[] persons, concerns or organizations listed below *

* NOTE: Separate verified statements are required from each person, concern or organization having rights to the invention averring to their status as small entities. (37 C.F.R. 1.27)

FULL NAME CANCER RESEARCH CAMPAIGN TECHNOLOGY LIMITED
ADDRESS CAMBRIDGE HOUSE, 6-10 CAMBRIDGE TERRACE, REGENT'S PARK, LONDON NW1 4JL

[] INDIVIDUAL [] SMALL BUSINESS CONCERN
[X] NON-PROFIT ORGANIZATION

FULL NAME _____
ADDRESS _____
[] INDIVIDUAL [] SMALL BUSINESS CONCERN
[] NON-PROFIT ORGANIZATION

FULL NAME _____
ADDRESS _____
[] INDIVIDUAL [] SMALL BUSINESS CONCERN
[] NON-PROFIT ORGANIZATION

I acknowledge the duty to file, in this application or patent, notification of any change in status resulting in the loss of entitlement to small entity status prior to paying, or at the time of paying, the earliest of the issue fee or any maintenance fee due after the date on which status as a small entity is no longer appropriate. (37 C.F.R. 1.28(b))

I hereby declare that all statements made herein on my own knowledge are true and that all statements made on information and belief are believed to be true; and further that these statements were made with the knowledge that willful false statements and the like so made are punishable by fine or imprisonment, or both, under Section 1001 of Title 18 of the United States Code, and that such willful false statements may jeopardize the validity of the application, any patent issuing thereon, or any patent to which this verified statement is directed.

NAME OF INVENTOR	NAME OF INVENTOR	NAME OF INVENTOR	NAME OF INVENTOR
GEOFFREY PAUL MARGISON	BRIAN MARPLES	SIMON SCOTT	JOLYON HINDSON HENDRY

Signature of Inventor	Signature of Inventor	Signature of Inventor	Signature of Inventor

Date	Date	Date	Date
14/6/01	01/10/01	11/10/01	14/6/01

Rec'd PCT/PTO 19 FEB 2002

HS

IN THE UNITED STATES PATENT AND TRADEMARK OFFICE

In re PATENT APPLICATION OF

BOX PCT

Geoffrey P. Margison et al.

Group Art Unit: Unassigned

Application No. 09/700,259

Examiner: Unassigned

Filed: November 14, 2000

Title: GENE THERAPY VECTORS AND THEIR USE IN ANTITUMOUR THERAPY

* * * * *

STATEMENT IN SUPPORT OF SEQUENCE LISTING

IN COMPLIANCE 37 C.F.R. §1.821 AND §1.825

Hon. Commissioner of Patents
Washington, D.C. 20231

Sir:

In accordance with 37 C.F.R. §1.821(f), Applicants' undersigned attorney hereby states that the content of the paper copy and the computer readable copy of the sequence listing submitted herewith are the same.

In accordance with 37 C.F.R. §1.821(g), Applicants' undersigned attorney hereby states that the submission herewith does not add any new matter to the application.

Respectfully submitted,

PILLSBURY WINTHROP LLP

By: Richard A. Steinberg
Richard A. Steinberg
Registration No. 26,588

1600 Tysons Boulevard
McLean, VA 22102
(703) 905-2000 Telephone
(703) 905-2500 Facsimile

Date: February 19, 2002

Attorney Reference: 040432/0275388

#3
RECORDED PTO 19 FEB 2002

SEQUENCE LISTING

<110> CANCER RESEARCH CAMPAIGN TECHNOLOGY LIMITED, et al

<120> IONIZING RADIATION OR DIATHERMY-SWITCHED GENE THERAPY
VECTORS AND THEIR USE IN ANTITUMOUR THERAPY

<130> PCT/GB99/01362

<140> PCT/GB99/01362

<141> 1999-05-17

<150> GB 9810423.5

<151> 1998-05-15

<160> 12

<170> PatentIn Ver. 2.1

<210> 1

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<223> Description of Artificial Sequence: Synthetic
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repeats of SEQ ID NO: 1.

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single stranded oligonucleotide sequence
complementary to SEQ ID NO: 4.

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for PCR amplification of Thymidine kinase gene.

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IN THE UNITED STATES PATENT AND TRADEMARK OFFICE

In re Application of

MARGISON ET AL

Serial No. 09/700,259

Group Art Unit: Unknown

Filed: November 14, 2000

Examiner: Unknown

For: GENE THERAPY VECTORS
AND THEIR USE IN ANTITUMOUR
THERAPY

October 30, 2001

AMENDMENT TO COMPLY WITH SEQUENCE LISTING RULES

Hon. Commissioner of
Patents and Trademarks
Washington, D.C. 20231

Sir:

In response to the Notification of Missing Requirements Under 35 U.S.C. § 371 dated April 30, 2001, Applicants submit the following amendment to comply with Sequence Listing Rules.

Amendments

Please delete the paper copy of the Sequence Listing filed on November 14, 2000. In its place, please enter the enclosed Sequence Listing on separate pages after the claims and abstract of the application.

Remarks

I. The Amendments

The specification of the application was amended to delete the Sequence Listing originally filed and to replace it with the substitute Sequence Listing enclosed herewith. The two Sequence Listings are substantively the same, but the enclosed Listing has been prepared using a more recent version of the program Patent-In. All of the sequence identification numbers referred to in the Sequence Listing were present in the application at the time of filing.

II. Submission of Computer Readable Copy of Sequence Listing

Applicants are including herewith a 3.5 inch computer readable diskette which contains a copy of the newly submitted Sequence Listing in ASCII text.

III. Statements to Comply With 37 C.F.R. §§ 1.821 and 1.825

In compliance with 37 C.F.R. § 1.821(f), Applicants' undersigned attorney hereby states that the content of the paper and computer readable copies of the Sequence Listing submitted herewith are the same. In accordance with 37 C.F.R. § 1.821(g), Applicants' undersigned attorney hereby states that the submission herewith does not add new matter to the application.

Conclusion

In light of the present amendments and enclosures, Applicants respectfully submit that all Sequence Listing requirements have now been complied with. It is

therefore respectfully submitted that this application is now in condition for substantive review.

If, in the opinion of the Examiner, a phone call may help to expedite the prosecution of this application, the Examiner is invited to call Applicants' undersigned attorney at (703) 905-2118.

Respectfully submitted,

PILLSBURY WINTHROP LLP


By _____
Paul N. Kokulis
Reg. No. 16773

PNK/MAS:ct
1600 Tysons Boulevard
McLean, Virginia 22102
Phone: (703) 905-2118

SEQUENCE LISTING

<110> Margison, Geoffrey

Marples, Brian

Scott, Simon

Hendry, Jolyon

<120> Gene Therapy Vectors and Their Use in Antitumor Therapy

<130> 40432/275388

<160> 12

<170> PatentIn version 3.0

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773
RECEIVED 30 OCT 2001

IN THE UNITED STATES PATENT AND TRADEMARK OFFICE

In re Application of

MARGISON ET AL

Serial No. 09/700,259

Group Art Unit: Unknown

Filed: November 14, 2000

Examiner: Unknown

For: GENE THERAPY VECTORS
AND THEIR USE IN ANTITUMOUR
THERAPY

October 30, 2001

PRELIMINARY AMENDMENT

Hon. Commissioner of
Patents and Trademarks
Washington, D.C. 20231

Sir:

Please amend the above application as follows:

IN THE CLAIMS

Cancel claims 51, 52 and 53.

Please enter the following amended claims:

3. (Amended) Vector material as claimed in Claim 2 wherein the tumour-sensitizing expression product is a prodrug activating enzyme selected from the group consisting of:

HSV thymidine kinase, nitroreductase, cytosine deaminase, cytochrome p450 2E1 and cytochrome p450 2DV1.

5. (Amended) Vector material as claimed in Claim 1 wherein the control gene encodes a recombinase enzyme that acts on recombinase target sites to modify the vector material to establish said operative linkage between the sensitizing gene expression regulatory system and the sensitizing gene or genes.

7. (Amended) Vector material as claimed in Claim 5 wherein said recombinase target sites are separated by a region containing a "stop" sequence of nucleotides that blocks or otherwise prevents expression of the sensitizing gene or genes until removed by the action of said recombinase enzyme.

8. (Amended) Vector material as claimed in Claim 5 wherein the protein coding regions of the sensitizing gene or genes are operationally separated from the said promoters and wherein said recombinase target sites are arranged such that recombination brings about the juxtapositioning of the sensitizing gene promoters and protein coding regions of the sensitizing gene or genes resulting in their expression.

9. (Amended) Vector material as claimed in Claim 5 wherein the control gene is a fusion gene that when expressed produces a fusion protein consisting of a recombinase and an intercellular trafficking protein.

10. (Amended) Vector material as claimed in Claim 5 wherein the region between said recombinase target sites contains a duplicate copy of the recombinase control gene together with an associated promoter.

11. (Amended) Vector material as claimed in Claim 1 wherein the sensitizing gene expression regulatory system incorporates at least one expression inducible element responsive to the effect of a predetermined exogenous or endogenous expression inducing influence.

12. (Amended) Vector material as claimed in Claim 1 wherein the sensitizing gene is a fusion gene that when expressed produces a fusion protein consisting of a sensitizing protein and an intercellular trafficking protein.

13. (Amended) Vector material as claimed in Claim 1 wherein the or each tumour sensitizing gene is selected from the group consisting of the *E. coli* nitroreductase gene, cytosine deaminase (CD) gene, *Herpes simplex* virus thymidine kinase (HSV-*tk*), mammalian cytochrome p450 2E1 or 2D1 gene, and their functional equivalents.

14. (Amended) Vector material as claimed in Claim 1 wherein the tumour cell sensitizing gene or genes and the control gene are in separate vectors.

15. (Amended) Vector material as claimed in Claim 1 wherein the tumour cell sensitizing gene or genes and the control gene are in the same vector.

16. (Amended) Vector material as claimed in Claim 1 for use in antitumour therapy, said use comprising the introduction of the vector material into tumour cells.

18. (Amended) Vector material as claimed in Claim 17 wherein at least one element of the control gene expression regulatory system is selected so that the control gene is automatically upregulated to an effective operational level when the vector material is introduced into cells of said tumours.

20. (Amended) Vector material as claimed in Claim 17 wherein said control gene expression regulatory system responds in use in a transfected cell to said expression inducing influence where the latter is provided by a change in environmental thermal conditions in cells containing the vector material.

21. (Amended) Vector material as claimed in Claim 17 wherein said control gene expression regulatory system responds in use in a transfected cell to said expression inducing influence where the latter is provided by a change in local oxygen concentration.

22. (Amended) Vector material as claimed in Claim 16 wherein said control gene expression regulatory system includes an expression control element responsive in use in a transfected cell to a hypoxia condition in the cellular environment.

25. (Amended) Vector material as claimed in Claim 23 wherein said control gene expression regulatory system responds in use in a transfected cell to said expression inducing agent which is provided by at least one of the following:

electromagnetic radiation, application of heat or cooling, application of a magnetic or electric field, an exogenous chemical inducing agent, radiation in the form of sub-atomic particles.

32. (Amended) Vector material as claimed in Claim 20 wherein the antitumour drug is selected from the group consisting of:

Temozolomide, Dacarbazine, Streptozotocin, Procarbazine, Carmustine, Semustine, Lomustine, Fotemustine, Busulphan, Treosulphane, Mechlorethamine, Cyclophosphamide, Iphosphamide, Chlorambucil, Melphalan, ethyleneimines, triethylene melamine, hexamethylmelamine, TEPA and thio-TEPA, dibromomannitol and dibromodulcitol, hydroxyurea, Methotrexate, azaserine Azathioprin, 5-azacytidine, 5-fluorouracil, cytosine arabinoside, 6-mercaptopurine, Allopurinol 6-thioguanine, deoxycoformycin, Tiazofurin, Acivicin, Pyrazofurin and p-aminolaevulinic acid, plant alkaloids such as Vinblastine, Vincristine and Vindesine, Etoposide and Teniposide, antitumour antibiotics such as Doxorubicin, Daunorubicin, Actinomycin, Bleomycins, Mytomycin, Mythramycin, Mitozantrone hormones such as oestrogen and progesterone.

33. (Amended) Vector material as claimed in Claim 23 wherein said expression inducing agent is an exogenous chemical inducing agent in the form of a hormone that interacts with a receptor molecule which interacts with a complementary hormone responsive element in the control gene expression regulatory system.

34. (Amended) Vector material as claimed in Claim 23 wherein the control gene expression regulatory system comprises a gene upregulation system that can be activated by a chemical agent.

35. (Amended) Vector material as claimed in Claim 22 containing a number of different control gene expression regulatory elements responsive to different expression inducing influences so as to be activated under a range of different conditions.

36. (Amended) Vector material as claimed in Claim 1 wherein at least one element of the sensitizing gene expression regulatory system is inducible in response to the effect of a predetermined exogenous or endogenous expression inducing influence.

38. (Amended) Vector material as claimed in Claim 1 wherein at least one element of the sensitizing gene expression regulatory system is selected for efficiency in the particular tumour(s) to which said antitumour therapy is directed, the selection being carried out using gene array technology.

39. (Amended) Vector material as claimed in Claim 1 which includes a plurality of tumour sensitizing genes providing a range of different expression products.

41. (Amended) A pharmaceutical composition comprising vector material as claimed in Claim 1 in association with a pharmaceutically acceptable carrier or excipient.

43. (Amended) A kit comprising one or more unit doses of vector material as defined in Claim 1 together with a transfection agent.

44. (Amended) A kit comprising:

- (a) a vector which comprises a tumour cell sensitizing gene or genes and a sensitizing gene expression regulatory system as defined in Claim 1;
- (b) a vector which comprises a control gene and a control gene expression regulatory system as defined in Claim 1; and
- (c) instructions for the use of vectors (a) and (b) in antitumour therapy.

45. (Amended) A kit comprising:

- (a) vector material as defined in Claim 1;
- (b) a vector which comprises a tumour cell sensitizing gene or genes and a sensitizing gene expression regulatory system as defined in Claim 1.

47. (Amended) A kit as claimed in Claim 43 wherein each of the vectors and/or vector material is provided in the form of a pharmaceutical composition in association with a pharmaceutically acceptable carrier or excipient.

48. (Amended) A kit as claimed in Claim 43 wherein the sensitizing gene(s) produce(s) a prodrug activating agent and said kit also includes at least one dose of a prodrug matched to said prodrug activating agent.

49. (Amended) A method of treatment for cancer patients wherein there is delivered to tumour cells vector material as claimed in Claim 1, said cells then being subjected to the appropriate expression inducing influence.

REMARKS

The claims are being amended to improve form.

Respectfully submitted,

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APPENDIX

Version with Markings to Show Changes Made

IN THE CLAIMS

Claims 51, 52 and 53 are being canceled.

The claims are amended as follows:

3. (Amended) Vector material as claimed in Claim 2 wherein the tumour-sensitizing expression product is a prodrug activating enzyme selected from the [following] group consisting of:

HSV thymidine kinase, nitroreductase, cytosine deaminase, cytochrome p450 2E1 [or] and cytochrome p450 2D1.

5. (Amended) Vector material as claimed in [any one of the preceding claims] Claim 1 wherein the control gene encodes a recombinase enzyme that acts on recombinase target sites to modify the vector material to establish said operative linkage between the sensitizing gene expression regulatory system and the sensitizing gene or genes.

7. (Amended) Vector material as claimed in Claim 5 [or 6] wherein said recombinase target sites are separated by a region containing a "stop" sequence of nucleotides that blocks or otherwise prevents expression of the sensitizing gene or genes until removed by the action of said recombinase enzyme.

8. (Amended) Vector material as claimed in Claim 5 [or 6] wherein the protein coding regions of the sensitizing gene or genes are operationally separated from the said promoters and wherein said recombinase target sites are arranged such that recombination brings about the juxtapositioning of the sensitizing gene promoters and protein coding regions of the sensitizing gene or genes resulting in their expression.

9. (Amended) Vector material as claimed in [any one of Claims] Claim 5 [to 8] wherein the control gene is a fusion gene that when expressed produces a fusion protein consisting of a recombinase and an intercellular trafficking protein [(such as for example the virion protein VP22)].

10. (Amended) Vector material as claimed in [any one of Claims] Claim 5 [to 9] wherein the region between said recombinase target sites contains a duplicate copy of the recombinase control gene together with an associated promoter.

11. (Amended) Vector material as claimed in [any one of the preceding claims] Claim 1 wherein the sensitizing gene expression regulatory system incorporates at least one expression inducible element responsive to the effect of a predetermined exogenous or endogenous expression inducing influence.

12. (Amended) Vector material as claimed in [any one of the preceding claims] Claim 1 wherein the sensitizing gene is a fusion gene that when expressed

produces a fusion protein consisting of a sensitizing protein and an intercellular trafficking protein.

13. (Amended) Vector material as claimed in [any one of Claims] Claim 1 [to 8] wherein the or each tumour sensitizing gene is selected from the group consisting of the *E. coli* nitroreductase gene, cytosine deaminase (CD) gene, *Herpes simplex* virus thymidine kinase (HSV-tk), mammalian cytochrome p450 2E1 or 2DV1 gene, and their functional equivalents.

14. (Amended) Vector material as claimed in [any one of the preceding claims] Claim 1 wherein the tumour cell sensitizing gene or genes and the control gene are in separate vectors.

15. (Amended) Vector material as claimed in [any one of Claims] Claim 1 [to 13] wherein the tumour cell sensitizing gene or genes and the control gene are in the same vector.

16. (Amended) Vector material as claimed in [any one of the preceding claims] Claim 1 for use in antitumour therapy [characterized in that the], said use comprises comprising the introduction of the vector material into tumour cells.

18. (Amended) Vector material as claimed in Claim 17 [further characterised in that] wherein at least one element of the control gene expression regulatory system is selected so that the control gene is automatically upregulated to

an effective operational level when the vector material is introduced into cells of said tumours.

20. (Amended) Vector material as claimed in [any one of Claims] Claim 17 [to 19] wherein said control gene expression regulatory system responds in use in a transfected cell to said expression inducing influence where the latter is provided by a change in environmental thermal conditions in cells containing the vector material.

21. (Amended) Vector material as claimed in [any one of Claims] Claim 17 [to 19] wherein said control gene expression regulatory system responds in use in a transfected cell to said expression inducing influence where the latter is provided by a change in local oxygen concentration.

22. (Amended) Vector material as claimed in [any one of Claims] Claim 16 [to 19] wherein said control gene expression regulatory system includes an expression control element responsive in use in a transfected cell to a hypoxia condition in the cellular environment.

25. (Amended) Vector material as claimed in Claim 23 [or 24] wherein said control gene expression regulatory system responds in use in a transfected cell to said expression inducing agent which is provided by at least one of the following:
electromagnetic radiation, application of heat or cooling, application of a magnetic or electric field, an exogenous chemical inducing agent, radiation in the form of sub-atomic particles.

32. (Amended) Vector material as claimed in Claim 20 wherein the antitumour drug is selected from the [following] group consisting of:

Temozolomide, Dacarbazine, Streptozotocin, Procarbazine, Carmustine, Semustine, Lomustine, Fotemustine, Busulphan, Treosulphane, Mechlorethamine, Cyclophosphamide, Iphosphamide, Chlorambucil, Melphalan, ethyleneimines, triethylene melamine, hexamethylmelamine, TEPA and thio-TEPA, dibromomannital and dibromodulcitol, hydroxyurea, Methotrexate, azaserine Azathioprin, 5-azacytidine, 5-fluorouracil, cytosine arabinoside, 6-mercaptopurine, Allopurinol 6-thioguanine, deoxycoformycin, Tiazofurin, Acivicin, Pyrazofurin and p-aminolaevulinic acid, plant alkaloids such as Vinblastine, Vincristine and Vindesine, Etoposide and Teniposide, antitumour antibiotics such as Doxorubicin, Daunorubicin, Actinomycin, Bleomycins, Mytomycin, Mythramycin, Mitozantrone hormones such as oestrogen and progesterone.

33. (Amended) Vector material as claimed in Claim 23 [or 24] wherein said expression inducing agent is an exogenous chemical inducing agent in the form of a hormone that interacts with a receptor molecule which interacts with a complementary hormone responsive element in the control gene expression regulatory system.

34. (Amended) Vector material as claimed in Claim 23 [or 24] wherein the control gene expression regulatory system comprises a gene upregulation system that can be activated by a chemical agent.

35. (Amended) Vector material as claimed in Claim 22 [or 23] containing a number of different control gene expression regulatory elements responsive to different expression inducing influences so as to be activated under a range of different conditions.

36. (Amended) Vector material as claimed in [any of the preceding claims]
Claim 1 wherein at least one element of the sensitizing gene expression regulatory system is inducible in response to the effect of a predetermined exogenous or endogenous expression inducing influence.

38. (Amended) Vector material as claimed in [any of the preceding claims]
Claim 1 wherein at least one element of the sensitizing gene expression regulatory system is selected for efficiency in the particular tumour(s) to which said antitumour therapy is directed, the selection being carried out using gene array technology.

39. (Amended) Vector material as claimed in [any of the preceding claims]
Claim 1 which includes a plurality of tumour sensitizing genes providing a range of different expression products.

41. (Amended) A pharmaceutical composition comprising vector material as claimed in [any of the preceding claims] Claim 1 in association with a pharmaceutically acceptable carrier or excipient.

43. (Amended) A kit comprising one or more unit doses of vector material as defined in [any one of Claims] Claim 1 [to 40] together with a transfection agent.

44. (Amended) A kit comprising:

- (d) a vector which comprises a tumour cell sensitizing gene or genes and a sensitizing gene expression regulatory system as defined in [any one of the Claims] Claim 1 [to 40];
- (e) a vector which comprises a control gene and a control gene expression regulatory system as defined in [any one of Claims] Claim 1 [to 40]; and
- (f) instructions for the use of vectors (a) and (b) in antitumour therapy.

45. (Amended) A kit comprising:

- (c) vector material as defined in [any one of Claims] Claim 1 [to 40];
- (d) a vector which comprises a tumour cell sensitizing gene or genes and a sensitizing gene expression regulatory system as defined in [any one of Claims] Claim 1 [to 40].

47. (Amended) A kit as claimed in Claim 43[, 44, 45 or 46] wherein each of the vectors and/or vector material is provided in the form of a pharmaceutical composition in association with a pharmaceutically acceptable carrier or excipient.

48. (Amended) A kit as claimed in [any one of Claims] Claim 43 [to 47] wherein the sensitizing gene(s) produce(s) a prodrug activating agent and [wherein one or more doses] said kit also includes at least one dose of a prodrug matched to said prodrug activating agent [are also included in said kit].

49. (Amended) A method of treatment for cancer patients wherein there is delivered to tumour cells vector material as claimed in [any of Claims] Claim 1 [to 40], said cells then being subjected to the appropriate expression inducing influence.

GENE THERAPY VECTORS AND THEIR USE IN ANTITUMOUR
THERAPY

FIELD OF THE INVENTION

The present invention relates to the field of molecular biology and
5 gene therapy, especially as applied to cancer therapy.

BACKGROUND

In connection with cancer therapy it has been suggested that cancer cells may be effectively treated by introducing into them sensitizing foreign genes, the expression of which leads to the destruction or elimination of these
10 cells. This could be achieved, for example, via the expression of a cytotoxic protein or cytotoxic RNA species, or via the expression of an immune-response stimulating factor or of substances that can bring about or promote the bio-conversion and activation of a systemically applied inactive chemical agent or prodrug to form an active cytotoxic drug (see for example Elizabeth A. Austin,
15 *et al.*, (1992), "A First Step in the Development of Gene Therapy for Colorectal Carcinoma: Cloning, Sequencing, and Expression of *Escherichia coli* Cytosine Deaminase", *Molecular Pharmacology*, 43, 380-387). The latter approach has been generally favoured in most cases because the active drug so produced is able to kill cells in the vicinity of the sensitised activating cells (the "bystander"
20 effect), thus compensating for any inefficiencies in cellular uptake of the activating gene expression system.

Difficulties with these gene therapy methods, however, include the fact that most current DNA or gene delivery and transfection systems usually propose the use of genetically engineered viruses which, up to now, have not
25 been able to deliver DNA for therapeutic purposes exclusively to tumour cells. Although there have been attempts to circumvent this problem by employing expression controlling regions (promoter and/or enhancer elements) of genes

that are predominantly expressed in tumour cells to direct the expression of the tumour-cell sensitizing mechanism, it has had to be accepted that such genes can also be expressed in normal tissues so that selectivity of expression is not absolute. Methods for the delivery of the genes specifically to target tumour 5 cells by exploiting cell-specific surface antibodies or receptors have also been considered, but these are also expressed in a number of normal cell types and thus selection is again not absolute. In order to be useful and safe for the treatment of cancer, ideally there needs to be a level or levels of selection that will result in expression of the cytotoxic mechanism in the tumour or in the 10 vicinity of the tumour or in the tumour cells exclusively.

Some methods of cancer treatment depend on ionizing radiation which is very commonly used against a variety of cancers. Devices are widely available for directing the radiation from an external source to the tumour in such a way that the dose of radiation to normal tissues is minimised (conformal 15 radiotherapy). However, problems associated with radiotherapy include the fact that tumour cells can often be more resistant to treatment than normal cells, whilst some normal cell types inside the radiation field may be very radiosensitive. In some alternative attempts to deliver the radiation more specifically only to tumour cells, tumour-targeting antibodies or similar 20 molecules have been labelled with various radioisotopes. In this latter case, however, the amounts of radiation that can be administered overall to the tumour within tolerable systemic levels are often too small to be of sufficient direct therapeutic value and this approach has therefore been generally considered more suitable for tumour imaging than for therapy.

25 The term "ionizing radiation" as used herein may include not only electromagnetic radiation such as X-rays and γ -rays, but also high energy sub-atomic particles such as α -particles and β -rays or electron beams.

Apart from chemotherapy, another method of cancer treatment uses diathermy. However, despite extensive trials, hitherto this method has not found widespread use because the response curves for human cells are very steep and because of difficulties in achieving and maintaining precise and 5 homogeneous elevated temperatures in deep-seated tumours.

One objective of the present invention is to provide improved means and methods for selectively killing or eliminating tumour cells using a low or transient dose of a gene expression inducing agent to switch on a gene that produces an expression product within tumour tissue that has the effect of 10 bringing about the destruction or removal of tumour cells.

SUMMARY OF THE INVENTION

The present invention relies at least in part on a concept of selectively bringing about the destruction or elimination of tumour cells by using gene therapy methods to provide the tumour cells with a silenced or dormant killing 15 mechanism that can be activated by exposing the cells to an appropriate stimulating influence, e.g. ionizing radiation, heat, or a chemical or other inducing agent, so as to upregulate a control gene which then triggers and switches on or primes a tumour cell sensitizing gene or genes. It is in fact already known that expression of certain genes in human cells can be induced 20 or upregulated by exposure of the cells to heat (e.g. from diathermic devices) or ionizing radiation, in the latter case sometimes after very low doses (see for example D.E. Hallahan *et al.* (1995) "Spatial and temporal control of gene therapy using ionizing radiation", *Nature Medicine*, Vol. 1, No. 8, 786-791) and D. E. Hallahan, *et al.*, (1995), "*c-jun* and *Egr-1* Participate in DNA Synthesis 25 and Cell Survival in Response to Ionizing Radiation Exposure", *The Journal of Biological Chemistry*, 270, 30303-30309), but it is believed that this effect has not previously been exploited in the same way as in the present invention.

More specifically, the invention envisages providing means for carrying out a method of treatment for cancer patients in which there are delivered to tumour cells vectors, preferably self-replicating viral or plasmid vectors, containing a dormant or silenced tumour cell sensitizing gene or genes 5 of which transcription and expression can be initiated by an inducible control mechanism responsive to a particular stimulatory influence which may be applied to the tumour cells concerned, or which may arise as an endogenous product of the tumour cells, whereby said sensitizing gene or genes are caused or enabled to generate a product that will bring about the death or elimination 10 of said tumour cells. e.g. through activation of a cytotoxic prodrug or other cytotoxic agent or tumour cell destruction or elimination mechanism.

Thus, from one aspect the invention provides vector material characterised in that contains:

- (a) a tumour cell sensitizing gene or genes of which expression in a 15 tumour cell yields a sensitizing gene expression product having a potential to cause tumour cells to be killed and destroyed, or to be eliminated, or otherwise to be inactivated, or to be rendered sensitive and/or vulnerable to destruction;
- (b) a sensitizing gene expression regulatory system, including 20 promoter means, for said sensitizing gene or genes;
- (c) at least one control gene; and
- (d) a control gene expression regulatory system responsive in use in a 25 transfected cell to the effect of a predetermined exogenous or endogenous expression inducing influence so as to induce expression of said control gene to yield an expression product having a capacity to establish an operative linkage between said sensitizing gene expression regulatory system and said sensitizing

gene or genes effective to trigger and switch on or permit continuous or permanent expression of the latter to bring about continuous production of said sensitizing gene expression product.

The term "gene" is used herein to denote one or more nucleotide sequences, with or without intervening introns, that encodes a functional protein or RNA molecule. It may therefore embrace cDNA sequences. Use of this term in the singular, and of other terms such as gene expression regulatory system, may also cover the plural, and vice versa, where the context so admits.

Also, the term "vector" is used herein to denote an agent or vehicle adapted to act as a carrier of nucleic acid fragments or nucleotide sequences inserted therein for the purpose of introducing such fragments or sequences into a prokaryotic (bacterial) or eukaryotic cell. As such, the term "vector" as used herein embraces viruses, including phages, and nucleic acid gene portions thereof, as well as bacterial and synthetic plasmids. Where the vector contains an inserted gene or genes and a regulatory system or promoter which facilitates efficient transcription and, where appropriate, translation of said inserted gene(s), it is termed an "expression vector".

The vector material of this invention is particularly suitable for use in antitumour therapy, in which case it will be introduced into tumour cells, and the following description will generally assume such use although this is not necessarily to be construed as a limitation to the scope of the invention.

In at least some cases where the control gene expression inducing agent is an exogenous agent, such as for example heat or ionizing radiation, it may be sufficient for this to be applied at sub-lethal doses, and/or at sub-therapeutic tumour imaging doses in the case of ionising radiation for example.

The invention may also be defined in one aspect as consisting of expression vector material for use in introducing into tumour cells in the course of antitumour therapy, characterised in that the said vector material contains a

tumour cell sensitizing gene (or genes) and also a gene expression control system adapted to be activated by a control gene expression inducing influence, such as for example ionizing radiation, a chemical inducing agent, non-ionizing electromagnetic radiation or diathermic heat, in such a way as to bring about or 5 permit permanent or continuous expression in transfected tumour cells of said tumour cell sensitizing gene(s), yielding an expression product that causes or enables said cells to be killed and destroyed, or to be eliminated, or otherwise inactivated, or to be rendered sensitive and/or vulnerable to destruction.

In general, the term "tumour cell sensitizing gene" is used herein to 10 denote a gene or any DNA sequence or combination thereof which when expressed *in vivo* in a tumour cell generates a product that is effective at least potentially in bringing about the destruction or elimination of such tumour cell and possibly other tumour cells in the vicinity or in metastases.

Various kinds of tumour cell sensitizing genes may be used to yield an 15 expression product that will give the desired result. In practice, it will often be preferred to use a tumour cell sensitizing gene (or combination of genes) which yields an expression product that is itself a cytotoxic agent or, often more preferably, that is an enzyme or other bioactive agent able to bring about the breakdown or conversion of an inactive prodrug into an active cytotoxic form 20 or otherwise promotes a cytotoxic effect of another potentially toxic agent. For example, the *Herpes simplex* virus thymidine kinase gene may be used to produce, when transcribed and translated, the thymidine kinase enzyme which is a prodrug activating agent able to convert the inactive prodrug gancyclovir into a cytotoxic metabolite. An effective amount of such prodrug may be 25 administered systemically, at the same time as transfected tumour cells are subjected to the appropriate control gene expression inducing influence, e.g. ionizing radiation, chemical agent or heat treatment, or before this treatment, or subsequent to this treatment. In another example a sensitizing gene may be

used which expresses an iodine transport protein which can have the effect of causing radioactive iodine, separately administered, to concentrate in target tumour cells such that the latter are killed.

Instead of or as well as providing tumour sensitizing genes or cDNAs that encode prodrug activating enzymes or other toxic agents, e.g. toxic proteins such as ricin, additional or alternative possibilities include (a) providing DNA sequences or cDNAs that encode immune response stimulating factors intended to bring about the elimination of not only the primary tumour cells but also other tumour cells in tumour metastases, (b) providing DNA sequences encoding ribozymes, oligoribonucleotides or RNA molecules, especially antisense molecules, that will attenuate the expression of vital proteins or RNA molecules, i.e. any such molecules that are essential for cell survival and propagation, and (c), providing any other cell killing or cell removal mechanisms that can be activated by the expression product of a sensitizing gene. Hereinafter, these various DNA sequences will be collectively referred to as "genes", and insofar as they may be administered to a mammal for therapeutic purposes they may all be regarded as covered by the term "therapeutic DNA".

Examples of immune response or cytokine genes that stimulate an immune response in the host in respect of tumour cells include genes that code for GM-CSF, IFN-alpha, IFN-beta, IFN-gamma, IL-1 β , IL-2, IL-4, IL-6, IL-8, IL-10, IL-12, IL-15 and TNF α . Other cell killing or sensitizing genes that may be used include Angiostatic genes such as angiostatin, endostatin, IP-10, Mig, PEX, Kringle-5, SDF-1 α , TIMP-1, TIMP-2, TIMP-3 and TIMP-4, apoptotic genes such as Bak, Bax, Bcl-XL, Bcl-XS, Bik, SARP-2 and TRAIL; cytolytic genes such as granzymeA, granzymeB and perforin; and gap junction genes such as Connexin26, Connexin32 and Connexin43.

The following points may be noted:

1. Angiostatic genes inhibit the formation of new blood vessels (caused via hypoxia) and would thus increase the level of hypoxia in the tumours, as well as killing (starving to death) cells in the region: (another bystander effect).
5 2. Apoptotic genes simply kill cells via apoptosis.
3. Cytolytic genes simply lyse cells.
4. Gap junction genes encode proteins (connexins) that are necessary for the bystander effect as mediated by activated prodrugs (see for example Elshami, A.A., *et al.* (1996) "Gap junctions play a role in the
10 'bystander effect' of the herpes simplex virus thymidine kinase/ganciclovir system in vitro", *Gene Ther.* 3, 85-92), so co-expression could enhance the effect.

In relation to the bystander effect, it may also be noted that bystander cell killing effects can be observed in tumours that have no direct contact with a
15 transduced tumour cell that expresses a prodrug activating gene ("distant" bystander effect; Wilson et al 1996; Kianmanesh et al 1997). This appears to be a consequence of an immune response mechanism (Caruso et al 1993; Kianmanesh et al 1997; Misawa et al 1997).

Although it is feasible to arrange for expression of tumour cell
20 sensitizing gene(s) to be directly under the control of an ionizing radiation or heat responsive promoter or enhancer or other inducible expression-regulatory element of the sensitizing gene expression regulatory system, a potentially serious practical problem results from the fact that such genes presently known which respond to give the highest levels of such inducing influences do so only
25 transiently. This suggests that in order to achieve effective treatment it would be necessary to apply continuous or repetitive exposure to heat or ionizing radiation or other expression inducing influence. Also, in the case in which

prodrugs are employed, it would be necessary to apply such prodrugs at a particular time for appropriate levels of prodrug activation to occur. Whilst continuous exposure, for example, to ionizing radiation at low doses sufficient for activating radiation responsive promoter elements may be achieved using a 5 suitable radioactive isotope-labelled tumour specific antibody or ligand, prima facie it would appear that any antitumour treatment methods based for example on exploiting the response of radiation responsive or heat responsive promoter elements to ionizing radiation or to heat or other exogenous expression inducing influence in a gene therapy scenario would necessarily be of limited 10 value. However, in embodiments of the present invention herein presented there is provided vector material so constructed that expression of the tumour cell sensitizing gene(s) is indirectly under the control of an expression inducing agent responsive element or elements of a control gene expression regulatory system, such element or elements being arranged to have the effect when 15 activated, albeit transiently, of triggering continuous expression of the tumour sensitizing gene(s), or at least of permitting or enabling continuous expression thereof or priming the cell transfected with such vector material for continuous expression of said sensitizing gene(s).

Continuous expression is achieved in accordance with the invention by 20 constructing the vector material so that activation of the control gene expression regulatory system induces expression of the control gene in the vector material which encodes an enzyme that acts as a trigger to modify the vector material, for example through a site-specific recombination system, in such a way as to switch on continuous expression of the tumour-sensitizing 25 gene(s), provided of course that the expression regulatory system of the tumour sensitizing gene(s) is also active. It will of course be appreciated that in some cases the recombination or other vector modification brought about by the expression product of the control gene may not necessarily result in immediate

continuous or permanent expression of the sensitizing gene(s) since the latter may also be conditional on the addition or presence of some other agent, e.g. a chemical agent such as a hormone, or a tetracycline or IPTG in a prokaryotic-based regulation system. In such cases the effect of the activation of the control
5 gene may be regarded as priming the tumour sensitizing gene(s), permitting expression of the latter.

Regarding the reference to prokaryotic-based expression regulation systems, such as for example the "Tet-On Tet-Off" regulation system and the isopropyl beta-D-thiogalactoside (IPTG) regulation system (see Gossen M,
10 Bonin AL, Freundlieb S, Bujard H "Inducible gene expression systems for higher eukaryotic cells", *Curr Opin Biotechnol* 1994 Oct;5(5):516-20), Gossen M, Freundlieb S, Bender G, Muller G, Hillen W, Bujard H "Transcriptional activation by tetracyclines in mammalian cells". *Science* 1995 Jun 23;268(5218):1766-9 and Wyborski DL, Short JM "Analysis of inducers of the
15 E.coli lac repressor system in mammalian cells and whole animals", *Nucleic Acids Res* 1991 Sep 11;19(17):4647-53), these are commercially available (e.g. Clontec and Stratagene) and could also be incorporated in the control gene expression regulatory system.

In some embodiments where antitumour therapy is involved the
20 expression inducing influence to which the control gene expression regulatory system of the vector material responds is endogenous and tumour related, being produced by tumour cells associated specifically with tumours to which said antitumour therapy is directed. In this case, at least one element of the control gene expression regulatory system may be selected so that the control gene is
25 automatically upregulated to an effective operational level when the vector material is introduced into cells of the tumours concerned. For example, in some embodiments the control gene expression regulatory system may respond in use in a transfected cell to an expression inducing influence provided by a

change in local oxygen concentration or, alternatively, by a change in environmental thermal conditions in cells containing the vector material, e.g. hyperthermia or hypothermia. To achieve high efficiency for a particular tumour type the selection of at least one element of the control gene expression regulatory system is conveniently carried out using gene array technology (see for example Schena-M; Shalon-D; Heller-R; Chai-A; Brown-PO; Davis-RW "Parallel human genome analysis: microarray-based expression monitoring of 1000 genes", *Proc-Natl-Acad-Sci-U-S-A.* 1996 Oct 1; 93(20): 10614-9, DeRisi-J; Penland-L; Brown-PO; Bittner-ML; Meltzer-PS; Ray-M; Chen-Y; Su-YA; 5 Trent-JM "Use of a cDNA microarray to analyse gene expression patterns in human cancer Nat-Genet", 1996 Dec; 14(4): 457-60, Duggan-DJ; Bittner-M; Chen-Y; Meltzer-P; Trent-JM "Expression profiling using cDNA microarrays". *Nat-Genet.* 1999 Jan; 21(1 Suppl): 10-4 and Watson-A; Mazumder-A; Stewart-M; Balasubramanian-S "Technology for microarray analysis of gene 10 expression", *Curr-Opin-Biotechnol.* 1998 Dec; 9(6): 609-14). Gene array technology may also be used to select efficient sensitizing gene regulatory elements for specific tumours.

In embodiments in which the expression inducing influence is provided by a change in local oxygen concentration this will generally result 20 from a condition involving a reduction in oxygen (hypoxia) within the tumour tissue. Many tumours normally contain regions of hypoxia, particularly where the blood supply has become inadequate owing to the overall proliferation of the tumour cells, so that the hypoxia is present endogenously. In addition, however, it is also known that an effect of ionizing radiation treatment of 25 human tumours is the transient induction of hypoxia within the tumours. These facts can be exploited in connection with the present invention since the promoters of many genes, e.g. the enolase-1 gene described by G.L. Semenza *et al* in *J. Biol. Chem.* 1996, 271, 32529-32537, contain elements responsive to

hypoxia inducing factor-1 (HIF-1) and expression of such genes has been shown to be induced by hypoxic conditions (see for example Wenger, R.H. and Gassmann, M. (1997) "Oxygen(es) and the hypoxia-inducible factor-1", *Biol Chem.* **378**, 609-616. Thus, to take advantage of this all that is necessary is to 5 incorporate a known hypoxia responsive element in the expression regulatory system or promoter of the control gene or genes in preparing the vector material of the present invention.

It would also be possible to synthesise a promoter that contains both radiation responsive and hypoxia-responsive elements so that Cre or other 10 recombinase enzyme would be expressed when either of the inducing conditions are fulfilled.

A hypoxia responsive element could also be included in the expression regulatory system or promoter of the tumour cell sensitizing gene or genes to control the expression of the sensitizing gene(s) if the tumour cell concerned 15 were to become hypoxic following priming of the cell after expression of the control gene and operation of the recombinase or other vector modification mechanism.

In most embodiments the expression inducing influence to which the control gene expression regulatory system responds in use in a transfected cell 20 is provided by an exogenous expression inducing agent applied to cells into which the vector material is introduced. Preferably, the dose level of such exogenous expression inducing agent needed to trigger and switch on expression of the sensitizing gene or genes in the cells of tissue into which the vector material has been introduced is substantially sub-lethal.

25 The exogenous control gene expression regulatory system or elements thereof may be selected to respond in use in a transfected cell to an expression inducing agent which will usually be provided by at least one of the following:

electromagnetic radiation, application of heat or cooling, application of a magnetic or electric field, an exogenous chemical inducing agent, radiation in the form of sub-atomic particles.

5 Where the expression inducing agent is electromagnetic radiation it may be in the form of ultra-violet or visible light, or ionizing radiation in the form of X-rays or gamma-rays. With regard to non-ionizing electromagnetic radiation, it is known for example that genotoxic stress produced by ultraviolet radiation can bring about expression of certain genes (see for example Liu ZG,

10 Baskaran R, Lea-Chou ET, Wood LD, Chen Y, Karin M, Wang JY "Three distinct signalling responses by murine fibroblasts to genotoxic stress", *Nature* 1996 Nov 21;384(6606):273-6), and visible light can also upregulate some genes such as the gene of heat shock protein 47 (HSP47) which responds to a wavelength of 652nm, the same wavelength as is used in some photodynamic

15 therapy (PDT) applications (see for example Nagata K Hsp47: "a collagen-specific molecular chaperone", *Trends Biochem Sci* 1996 Jan;21(1):22-6, Verrico AK, Moore JV "Expression of the collagen-related heat shock protein HSP47 in fibroblasts treated with hyperthermia or photodynamic therapy", *Br J Cancer* 1997;76(6):719-24 and Gomer CJ, Ryter SW, Ferrario A, Rucker N,

20 Wong S, Fisher AM "Photodynamic therapy-mediated oxidative stress can induce expression of heat shock proteins". *Cancer Res* 1996 May 15;56(10):2355-60). Radiation in the form of sub-atomic particles may be provided by radioactive isotopes. The effect of magnetic fields in inducing gene expression or upregulation has been reported by Goodman R, "Blank M

25 Magnetic field stress induces expression of hsp70", *Cell Stress Chaperones* 1998 Jun;3(2):79-88.

In some embodiments ionizing radiation in the form of α -particles may be generated *in situ* in tumour cells transfected with the vector material of this

invention by arranging for the cells to take up a boron compound and then irradiating with low energy neutrons. Since boron compounds are preferentially retained in certain tumour tissue, e.g. in the brain, this so-called boron neutron capture therapy (BNCT) technique (explained more fully in U.S. Patent No. 5599796 of Schinazi *et al.* of which the content is incorporated herein by reference) can provide a useful method of achieving selective antitumour treatment.

Where the expression inducing agent is an exogenous chemical inducing agent it may be a chemical agent that produces cellular damage, e.g. DNA damage or cell membrane damage or other oxidative damage. Such exogenous chemical inducing agent may in the form of an antitumour drug, e.g. a platinum containing drug such as cis-diaminedichloroplatinum (CDDP), commonly known as cisplatin. In this respect it has in fact been found that many expression regulatory elements that are responsive to ionizing radiation are also sensitive to platinum-containing antitumour drugs.

In some cases the expression inducing agent may be an exogenous chemical inducing agent in the form of a hormone that interacts with a receptor molecule which interacts with a complementary hormone responsive element in the control gene expression regulatory system. For examples of hormone control of the expression of specific genes via the interaction of the hormone with a hormone receptor and its subsequent binding, along with accessory factors to the promote regions of hormone responsive genes, see Ferlini C, *et al.* "Tamoxifen induces oxidative stress and apoptosis in oestrogen receptor-negative human cancer cell lines", *Br J Cancer* 1999 Jan;79(2):257-63 and Pratt MA, Satkunaratnam A, Novosad DM "Estrogen activates raf-1 kinase and induces expression of Egr-1 in MCF-7 breast cancer cells", *Mol Cell Biochem* 1998 Dec;189(1-2):119-25.

In practice there is likely to be a wide range of chemotherapeutic

agents or drugs that can be used in combination with appropriate promoter elements as expression inducing agents, especially drugs that have the effect of directly, or indirectly through cellular damage and/or signal transduction pathways, upregulating the expression of endogenous or exogenous genes.

5 Potential expression inducing drugs include those that directly or indirectly induce DNA damage such as the alkylating agents. These include methylating agents such as Temozolomide, Dacarbazine, Streptozotocin, and Procarbazine, the nitrosoureas such as Carmustine, Semustine, Lomustine, and Fotemustine, the alkyl sulphonates such as Busulphan and Treosulphan, the nitrogen mustards such as Mechlorethamine, Cyclophosphamide, Iphosphamide, Chlorambucil and Melphalan, the ethyleneimines such as triethylene melamine, hexamethylmelamine, TEPA and thio-TEPA, the epoxides such as dibromomannital and dibromodulcitol, the antimetabolites such as hydroxyurea, Methotrexate, azaserine, Azathioprin, 5-azacytidine, 5-10 fluorouracil, cytosine arabinoside, 6-mercaptopurine, Allopurinol, 6-thioguanine, deoxycoformycin, Tiazofurin, Acivicin, Pyrazofurin and p-aminolaevulinic acid, plant alkaloids such as Vinblastine, Vincristine and Vindesine, Etoposide and Teniposide, antitumour antibiotics such as Doxorubicin, Daunorubicin, Actinomycin, Bleomycins, Mytomycin, 15 Mythramycin, Mitozantrone, hormones such as oestrogen and progesterone, and analogues of these agents. In some circumstances, even aspirin may act as a chemical inducing agent, as reported by Fawcett TW, Xu Q, Holbrook NJ "Potentiation of heat stress-induced hsp70 expression in vivo by aspirin". *Cell Stress Chaperones* 1997 Jun;2(2):104-9.

20 It will accordingly be appreciated that the control gene expression regulatory system (and possibly also the tumour sensitizing gene expression regulatory system) can comprise a gene upregulation system that can be activated by a wide range of various chemical agents.

It can be noted that some further information about prodrug activating genes (including fusion genes) and also a list of immune response modifying genes is to be found, *inter alia*, on a web site www.invivgen.com which is to be regarded as part of this disclosure.

5 To provide tumour targeting the inducing agents used may be incorporated in liposomes or may be conjugated to other tumour targeting molecules.

In preferred embodiments the control gene encodes a recombinase enzyme that acts on recombinase target sites to modify the vector material to 10 establish the required operative linkage between the sensitizing gene expression regulatory system and the sensitizing gene or genes. Preferably the control gene and the recombinase target sites are part of a Cre-loxP or a Flp-frt site specific recombination system.

In these embodiments, the recombinase target sites are separated by a 15 region containing a "stop" sequence of nucleotides that blocks or prevents expression of the sensitizing gene or genes until removed by the action of said recombinase enzyme.

Thus, in some preferred embodiments the vector material has an inducible promoter system, including for example an ionizing radiation or heat 20 responsive promoter element or elements, operatively linked to a control gene that encodes a recombinase enzyme that is able to bring about the recombination of individual short segments of specific DNA sequences in another region of the same vector, or in a separate cotransfected vector, which in turn brings about a resultant activation and expression, or priming for 25 expression, of the tumour cell killing or sensitizing gene(s). Such specific DNA sequences constitute recombinase target sites and preferably they flank a region called a "Stop cassette" (available for example in Gibco™ plasmid pBS302), which contains transcription termination or stop sequences or which

contain some other intervening sequence that prevents expression of the downstream sequences (n.b. Gibco and Gibco BRL are trade marks of Gibco Europe Limited and/or Life Technologies, Inc.). Such "Stop cassette" is located upstream of the tumour sensitizing gene(s) but downstream of a separate promoter sequence or sensitizing gene expression regulatory system whereby expression of this tumour sensitizing gene is normally prevented. In these embodiments the vector is constructed so that the recombination of the DNA sequences of the recombinase target sites results in the elimination or deletion of this Stop cassette which is excised from the vector so that transcription and expression of the tumour killing or sensitizing gene(s) is then no longer prevented.

It will be understood that the control gene expression regulatory system, i.e. the promoter or enhancer element or elements thereof, will generally be such that exposure to the appropriate expression inducing agent, e.g. heat or ionizing radiation, elicits a response that brings about a substantial, or at least effective, increase in activity and hence transcription and translation of downstream sequences. Preferably the expression control or regulatory system is such that this response can be brought about by very low, non-therapeutic, sublethal doses of the expression inducing agent. Following activation by such treatment, the promoter or other responsive element(s) will then cause expression of the gene encoding the recombinase enzyme that brings about the sequence-specific recombination of the recombinase target sites which are located in another region of the same vector or in another vector cotransfected therewith, these target sites flanking the region referred to as the "Stop cassette".

Suitable recombinase genes that may be used in this arrangement include the *Escherichia coli* P1 Bacteriophage *cre* and the *Saccharomyces*

cerevisiae flp recombinase genes. Other genes with similar characteristics could also be used.

It should be pointed out that as the gene expression regulatory or control system in eukaryotic cells may comprise a relatively complex promoter consisting of a number of different, non-contiguous, separate parts, and may also include a more remotely located enhancer sequence associated with the promoter, the term promoter element or elements used herein is to be construed broadly as denoting any appropriate part of the expression regulatory or control system.

The *cre* gene, which expresses the Cre recombinase protein of bacteriophage P1, is used in conjunction with *loxP* target or recombination sites. The Cre-loxP site-specific recombination system is a well known recombination system (see for example Martina Anton. *et al.*, (1995), "Site-Specific Recombination Mediated by an Adenovirus Vector Expressing the Cre Recombinase Protein: a Molecular Switch for Control of Gene Expression", *Journal of Virology*, 69, 4600-4606, and Minmin Qin. *et al.*, (1995), "Site-specific cleavage of chromosomes in vitro through Cre-Lox recombination", *Nucleic Acids Research*, 23, 1923-1927), and the elements thereof, including the *cre* gene, *loxP* sites and Stop cassette assembled in plasmid vectors, are commercially available, e.g. as a Gibco BRL™ product from Life Technologies. Inc. (U.S.A.). The Flp site specific recombinase system in conjunction with FRT target sites that provide substrates for the Flp recombinase protein (see for example Dymecki (1996) "Flp recombinase promotes site-specific DNA recombination in embryonic stem cells and transgenic mice", *Proc. Natl. Acad. Sci. U.S.A.*, 93, 6191-6196; also U.S. Patents 5654182 and 5677177 of Wahl *et al.*) may be applied in a similar way to excise a blocking Stop cassette so as to "switch on" expression of the tumour

sensitizing gene after activation of the *fhp* recombinase gene which is arranged to be under the control of a radiation or heat responsive promoter in the vector.

Examples of promoters or expression control elements that can be activated by low doses of ionizing radiation include the enhancers and/or 5 promoters or expression control radiation responsive elements of the *egr-1* gene, TNF α gene, the *NfkB* gene, the *c-fos* gene, the *jun-b* gene, the *c-jun* and the *c-myc* gene. This list, however, is not exhaustive. A typical example of a heat responsive promoter that may be used is the *hsp-90* gene.

As indicated above, transcription and translation of the tumour 10 sensitizing gene(s) in the vector generally will be under the control of a separate sensitizing gene promoter or expression regulatory system. This is or includes preferably a promoter that operates very effectively in human cells and most preferably one that will operate specifically in the type of cells that make up the tumour to be treated, by virtue for example of being tissue or cell-type 15 specific or being associated with a tissue specific enhancer region, or even more preferably one that operates only in tumour cells. Conveniently, this promoter or regulatory system for activating the tumour sensitizing gene(s) may also be selected for good efficiency by the use of gene array technology as previously mentioned and should be located in the vector in such a position that 20 it does not bring about the expression of any gene at the protein level until a recombination of the DNA sequences of the recombinase target sites occurs. Thus, this sensitizing gene promoter or regulatory system may be located upstream of the Stop cassette which, as pointed out above, is itself located upstream of the tumour-sensitizing gene(s) so as to block or prevent expression 25 of the latter. Alternatively, this sensitizing gene promoter or regulatory system may be located within a region excised by the action of the recombinase enzyme such that upon recombination it is reorientated so as to be able to

promote continuous transcription and translation of said tumour-sensitizing gene(s) co-excised therewith.

In some embodiments the region between the recombinase target sites contains a duplicate copy of the recombinase control gene together with an associated promoter, this being one example of possible embodiments in which there is more than one control gene system allowing, for example, permanent expression of a recombinase gene or genes after the initial activation by the primary inducing agent.

Examples of mammalian cell promoters for the tumour sensitizing gene(s) that will generally be suitable include the human cytomegalovirus (CMV) gene promoter and the chicken B-actin gene promoter, whilst one example of a tissue specific promoter (in this case appropriate for treating prostatic tumours) is the prostate specific antigen promoter and its associated enhancer region (see for example E. R. Schuur, *et al.*, (1996), "Prostate-specific Antigen Expression Is Regulated by an Upstream Enhancer", *The Journal of Biological Chemistry*, 271, 7043-7051). Examples of tumour-specific promoters, in these cases acting in tumour cells that have lost p53 function, include the HSP-70 promoter (Tsutsumi-Ishii *et al.*, (1995) *Cell growth and differentiation*, 6, 1-8) and the MDR-1 promoter (Zastawy, R.L. (1993) *Oncogene* 8, 1529-1535). For tumour cells that have lost RB-1 function the E2F-1 promoter would be appropriate (M.J. Parr *et al.*, (1997) (see "Tumour-selective transgene expression in vivo mediated by an E2F-responsive adenoviral vector", *Nature Medicine*, 3, 1145-1149).

It will be understood that the sensitizing gene expression regulatory system may incorporate at least one expression inducible element responsive to the effect of a predetermined exogenous or endogenous expression inducing influence in a manner similar to the control gene expression control regulatory system.

It will of course be appreciated that although the control gene expression product will generally act to establish an operative linkage between the sensitizing gene expression regulatory system and the sensitizing gene or genes so as to permit expression of the latter, whether or not expression occurs 5 or continues permanently may depend on the presence of another agent needed to activate the promoter or other expression regulatory element of said sensitizing gene(s). Thus, in the absence of such other agent the action of the control gene expression product may be in effect to switch the sensitizing gene(s) into a "primed" mode ready to yield the active expression product when 10 the sensitizing gene expression regulatory system or promoter is activated. Statements in the present specification should be construed accordingly.

The vector material will generally be composed of vectors constructed in a known manner to provide, as necessary for optimal effect, a polyadenylation signal encoding region at the downstream ends of the 15 recombinase and tumour sensitizing gene(s), and appropriate eukaryotic cell origins of replication. For amplification in bacteria, there will also be included a bacterial origin of replication and an antibiotic resistance gene, as understood by those versed in the art.

As already mentioned, preferred tumour-sensitizing gene(s) include 20 genes that encode prodrug activating enzymes, but other examples are genes that encode cytotoxic proteins or toxins, immune response stimulating factors, ribozymes or antisense RNA molecules. Apart from the *Herpes simplex* virus (HSV) thymidine kinase (*tk*) gene previously mentioned, further examples of tumour sensitizing prodrug activating genes that can be suitable include the 25 *E.coli* nitroreductase (*nr*) gene (see S. M. Bailey *et al.*, (1996) "Investigation of alternative prodrugs for use with *E. coli* nitroreductase in 'suicide gene' approaches to cancer therapy" *Gene Therapy*, 12, 1143/1150), the cytosine deaminase gene, and the mammalian cytochrome p450 2E1 or 2DVI genes.

Other genes that encode proteins involved in different cell killing mechanisms can also be useful.

In a further development, the control gene is a fusion gene that when expressed also produces a fusion protein consisting of an intercellular trafficking protein (such as for example the virion protein VP22 – see for example Phelan A, Elliott G, O'Hare P “Intercellular delivery of functional p53 by the herpesvirus protein VP22”, *Nat Biotechnol* 1998 May;16(5):440-3 and Elliott G, O'Hare P Intercellular trafficking and protein delivery by a herpesvirus structural protein. *Cell* 1997 Jan 24:88(2):223-33) and a recombinase protein. Also, the tumour cell sensitizing gene or genes may comprise a fusion gene sequence that when expressed produces an intercellular trafficking protein and a tumour cell sensitizing protein.

In some cases, the tumour cell sensitizing gene or genes and the control gene are in separate vectors, but in other cases the tumour cell sensitizing gene or genes and the control gene are in the same vector.

If desired, it is also possible to include in the vector material a number of different control gene expression regulatory elements responsive to different expression inducing influences so as to be activated under a range of different conditions.

Also, in some embodiments a plurality of tumour sensitizing genes will be incorporated which may provide a range of different expression products for killing or eliminating the tumour cells. This may be useful, for example, where the sensitizing genes are intended to express prodrug activating agents – by arranging for a set of sensitizing genes to produce a range of different prodrug activating agents the most efficient prodrug may be selected for systemic administration in each particular case of patient treatment.

It is also contemplated that the vector material of this invention may be presented as a kit which may comprise one or more unit doses of the vector material, herein defined, together with a transfection agent. In such kits the control gene(s) and sensitizing gene(s) may be contained either in the same vectors or in separate vectors, and where the sensitizing gene(s) are designed to express prodrug activating agents for use in antitumour therapy, the kits may also include pre-prepared doses of a pre-selected prodrug or a range or prodrugs to be selected at the time of administration. The kits may also contain a range of different transfection agents that also may be selected at the time of administration.

The provision of such kits constitutes a further aspect of the invention.

The vector for delivery of the therapeutic DNA to patients can be a retroviral, lentiviral, adenoviral, adenovirus-associated viral, or an Epstein-Barr viral based vector or any viral or bacterial vector delivery system that might be used for gene therapy in humans. Alternatively, it can be a non-viral vector that would be made up for administration in a suitable formulation such as, for example, a complex with cationic liposomes or with a tumour-targeting antibody or ligand, or that would be incorporated into some other non-viral DNA delivery system for delivery to tumour tissue, especially human tumour tissue. In general, the vector will have the ability to invade tumour cells and to express the encoded tumour sensitizing genes(s) therein following exposure of transfected cells to the appropriate inducing agent. Delivery of the vector material will usually be carried out according to generally accepted gene therapy procedures or methods as described, for example, by Friedman in *Therapy for Genetic Disease*, T. Friedman, ed., Oxford University Press (1991) and reviewed by I.M.Verma and N. Somia (1997) in "Gene therapy-promises, problems and prospects" *Nature* 389, 239-242. For example, after constructing the viral or plasmid vector or vectors containing the tumour cell sensitizing

gene(s) and expression control element(s) described, this vector material may be incorporated in a pharmaceutical composition, possibly in combination with a pharmaceutically acceptable excipient or carrier vehicle and a transfection agent, for example a transmembrane carrier such as Penetratin™. This 5 composition may then be injected into the patient, either locally at the site of the tumour or systemically. Such pharmaceutical compositions or formulations represent a further aspect of the invention.

As indicated above, gene transfer methods known in the art which may be useful in the practical application of the present invention include both viral 10 and non-viral transfer methods. Viruses that have been used as gene transfer vectors include for example papovaviruses, vaccinia viruses, herpesviruses, as well as adenoviruses, lentiviruses, adeno-associated viruses and retroviruses of avian, murine and human origin. Many human gene therapy protocols have been based on disabled murine retroviruses.

15 Non-viral DNA transfer methods known in the art include mechanical techniques such as micro-injection, membrane fusion-mediated transfer via liposomes (as already mentioned), and direct DNA uptake and receptor-mediated DNA transfer. Also, viral-mediated DNA transfer can be combined with direct *in vivo* DNA transfer using liposome delivery which may allow one 20 to direct the viral vectors to the tumour cells concerned rather than into the surrounding normal cells. Alternatively, one may inject into tumours a self-replicating retroviral vector producer cell line so that there could then be a continuous source of DNA vector particles, similar to a technique already approved for use in humans with inoperable brain tumours. Although the 25 vectors may be taken up directly by cells, actively or by diffusion, liposome mediated transfer may be best achieved in some cases by use of a transfection agent such as a cationic lipid, e.g. the compound N-[1-(2,3-dioleoyloxy)-propyl]-N,N,N-trimethylammonium methylsulfate, commonly known as

DOTAP, marketed by Boehringer Mannheim. However, many other suitable methods will be known to persons familiar with gene therapy techniques. An important aspect of many embodiments of the invention is that the vector delivery system may not need to have absolute specificity for tumour cells since

5 the tumour cell killing mechanism will only be activated in the area affected by the expression inducing agent which may be accurately targeted, e.g. as with radiation or diathermy, and in some embodiments the killing mechanism may only operate in cancer cells by virtue of tumour-specific promoters driving the expression of the tumour-sensitizing gene(s).

10 For clinical use, the vectors may be mixed with a selected transfection agent to provide a pharmaceutical preparation which may be administered by any suitable means, for example parenterally, orally or perhaps topically. In at least some cases such pharmaceutical preparations will be in the form of a sterile liquid formulation, presented possibly in unit dosage form in sealed

15 ampoules ready for use, and as already mentioned delivery or administration may be effected by injection, e.g. directly into tumour tissue or intravenously. In practice, not only the method of administration but also the particular protocol employed may be important; however, the precise details of the treatment and appropriate dosages will generally be determined by carrying out

20 straightforward trials and by the general experience of the medical practitioners in charge of the treatment.

The invention also extends to methods for treating tumour cells in a biological host system which, at least in one embodiment, comprises:

(a) administering to the system an effective amount of a composition

25 comprising vector material containing a tumour cell sensitizing gene or genes and having a control gene expression regulatory system responsive to a predetermined exogenous or endogenous expression inducing influence, said control gene expression regulatory system

being operatively linked to a recombinase gene, together with recombinase target sites flanking a region of which removal permits continuous expression of said tumour cell sensitizing gene(s) as specified above, or primes said sensitizing gene(s) for continuous expression;

5 (b) causing said tumour cells transfected with said vector material to be subjected to a dose of an expression inducing agent effective to activate the recombinase gene expression control system of said vector material, thereby to bring about, via recombinase-mediated site specific recombination within the vector material, expression of the or each tumour cell sensitizing gene component;

10 and, in the case of prodrug activating genes,

15 (c) administering to the host an effective amount of a composition comprising a prodrug convertible into an active form by the expression product of said tumour cell sensitizing gene or genes.

Usually, in most embodiments, the dose of the expression inducing agent, e.g. diathermy heat or ionizing radiation, will be derived from a directed external source or from a radioisotopically labelled tumour-seeking or cell or tissue-type seeking agent. Examples of suitable radioisotopically labelled 20 tumour-seeking agents for use in the above method include materials such as metaiodobenzylguanidine (MIBG) or suitably radioisotopically labelled tumour cell specific antibodies or ligands.

The prodrug referred to above may for example comprise any of the following compounds, administered either singly or in appropriate 25 combinations: gancyclovir, CB1954, fluorouracil, dacarbazine, or cyclophosphamide, subject to the prodrug activating gene(s) being, respectively, a gene or genes or cDNAs encoding HSV thymidine kinase,

nitroreductase, cytosine deaminase, cytochrome p450 2E1 or cytochrome p450 2DVI.

To summarise, in general in preferred embodiments:

- (1) where an ionizing radiation or platinum antitumour drug responsive expression control system is used, the responsive element or elements is or are selected from the radiation responsive enhancer or promoter of the *egr-1* gene, the *TNF α* gene, the *Nfk β* gene, the *c-fos* gene, the *jun-b* gene, the *c-jun* the *c-myc* gene and their functional equivalents, and either a single such radiation responsive element may be used or a plurality of tandemly arranged radiation-responsive DNA sequence elements (all the same or different) may be used in an array thereof;
- (2) where a heat-responsive promoter or expression control element is used, this is the promoter of the *hsp-90* gene or its functional equivalent;
- (3) the recombinase gene contained in the vector material will be the *E.coli* bacteriophage P1 *cre* gene or the *Saccharomyces cerevisiae FLP* gene, or a functional equivalent of one or other of these genes;
- (4) the DNA sequence of the vector material will include polyadenylation signal encoding regions from the virus SV40 or equivalent inserted at appropriate locations;
- (5) the vector material may contain a mammalian cell promoter which is that of the cytomegalovirus (CMV) or that of the chicken B-actin gene, or a functional equivalent of these;
- (6) the vector material may contain a tissue specific promoter (e.g. the prostate specific antigen promoter and its associated enhancer region, or an equivalent of this) that is specific for the particular tumour type to be treated;

(7) the vector material may contain a tumour specific promoter such as the HSP-70 promoter, the MDR-1 promoter, the E2F-1 promoter or telomerase-related promoters (or their equivalents), according to the p53 or RB-1 or telomerase status of the tumour;

5 (8) the DNA vector material will contain recombinase target sites provided by the *loxP* sites (or equivalents) that are substrates for the Cre recombinase enzyme, or recombinase target sites provided by FRT sites (or their equivalents) that are substrates for the Flp recombinase enzyme, or equivalents;

10 (9) the vector material will contain a Stop cassette provided by a sequence that prevents expression of any protein coding sequence located downstream thereof unless this region is excised by the action of the recombinase, or by another mechanism that results in a change in orientation or loss of this Stop sequence;

15 (10) the tumour sensitizing gene is selected from the *E.coli* nitroreductase gene, cytosine deaminase (CD) gene, *Herpes simplex* virus thymidine kinase (HSV-*tk*), mammalian cytochrome p450 2E1 or 2DVI gene, and their functional equivalents;

20 (11) The vector or each vector of the vector material will contain a bacterial origin of replication;

(12) The vector or each vector of the vector material will contain at least one mammalian cell origin of replication;

(13) The vector or each vector of the vector material will contain a bacterial antibiotic resistance gene;

25 (14) The vector or each vector of the vector material will contain intron sequences;

(15) The vector or each vector of the vector material may contain a mammalian cell antibiotic resistance gene.

The invention also includes use of vector material or vectors as
5 hereinbefore specified in the manufacture of a medicament or of a kit for use in
antitumour therapy.

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By way of example of the manner in which the invention may be carried out illustrative embodiments and background work in developing the invention will now be described in more detail with reference to the accompanying drawings. The particular embodiments and examples illustrated and described, however, should not be construed in any way as a limitation on the scope of the invention.

BRIEF DESCRIPTION OF THE DRAWINGS

Schematic diagrams (not to scale) of vectors in accordance with the invention and containing the above-mentioned elements are shown in the accompanying drawings. In these drawings:

FIGURE 1 represents one embodiment of a gene therapy plasmid vector, herein labelled "pComplete1", containing an ionizing radiation responsive recombinase expression control system;

FIGURE 2 shows the structure and some of the components making up pComplete2, a modified version of the vector in Figure 1 (the curved arrows adjacent to the Cre and tk ORFs indicate the direction of Cre and tk transcription following recombination at the loxP sites);

FIGURE 3 shows the structure and some of the components making up pComplete3, another version of the vector illustrated in Figure 1;

FIGURE 4 is a diagram showing different stages in the construction of the pComplete1 vector which contains tandemly arranged radiation-responsive elements of the *egr-1* promoter (double-framed boxes indicate starting materials);

FIGURE 5 is a schematic diagram demonstrating how pComplete1 responds to ionising radiation, undergoes recombination, and permanently expresses the tumour sensitizing gene thymidine kinase resulting in gancyclovir activation and cell killing;

FIGURE 6 is a diagram showing stages in the construction of vectors pEGRL(b)-*cre* and pStop-*gfp* which were introduced into MCF-7 cells and used in testing the principle of the invention;

FIGURE 7 is a bar chart diagram showing FACS analysis of MCF-7
5 cells following transfection with pStop-*gfp* only (light grey bars) or pStop-*gfp* together with pEGRL(b)-*cre* (dark grey bars) and exposure to 5Gy or 10Gy (in two 5Gy doses) of ionising radiation. Controls were not irradiated (0Gy);

FIGURE 8 is a diagram showing the results of a further test in which
10 MCF-7 cells were transfected with pEGRL(b)-*cre* and selected using the antibiotic G418, giving rise to the cell clone pCE which was, in turn, transfected with pStop-*tk* and then subjected to radiation. in the absence (open circles) of and in the presence (closed circles) of the prodrug ganciclovir;

FIGURE 9a and 9b are diagrams showing tumour cell growth *in vitro*
in the presence of the prodrug ganciclovir, after increasing doses of radiation.
15 Figure 9a relates to MCF-7/E4*cre* cells (○), MCF-7/E4*cre* cells transfected with plasmid p*Stk* (●) and MCF-7 cells transfected with p*tk* (▼). Figure 9b MCF-7 cells (○), MCF-7 cells transfected with pE4*tk* (■). Error bars are shown in one direction only for clarity;

FIGURE 10 is a bar chart diagram illustrating induction of
20 fluorescence in MCF-7 cells transfected with plasmid vectors pE4GFP and exposed to cisplatin (1 micromolar), ionising radiation (5Gy) or both. Controls were unexposed; and

FIGURE 11 is a bar chart diagram illustrating FACS analysis results
for induction of fluorescence in NB1G human neuroblastoma cells transfected
25 respectively with plasmids pEGFP (which contains a hypoxia-responsive promoter) after incubation in the absence or presence of 25 micromolar CoCl₂ in the cell culture medium.

DETAILED DESCRIPTION OF SPECIFIC EMBODIMENTS

Referring to the drawings, in the particular example of the plasmid vector pComplete1 depicted in Figure 1 the ionizing radiation responsive recombinase expression control system is provided by a promoter sequence comprising a synthetic tandem array of radiation-responsive elements corresponding to radiation-responsive elements found in the promoter of the *egr-1* gene, and is located upstream of the cytomegalovirus immediate-early (CMV I.E.) promoter. A control gene providing a recombinase protein coding sequence or open reading frame (ORF), which in this example is that of the P1 bacteriophage *cre* gene (labelled *Cre* ORF), is located downstream of an intron sequence, the latter serving to enhance the translation of the downstream ORF. There is also in this plasmid vector a strong constitutive mammalian cell enhancer/promoter which is that of the cytomegalovirus gene (labelled CMV I.E. Enhancer/promoter), *loxP* recombinase target sites that provide substrates for the Cre recombinase enzyme, and a tumour sensitizing gene which in this example is the Herpes simplex virus thymidine kinase (*tk*) gene (labelled *tk* ORF). As indicated, the *loxP* sites flank a Stop cassette. Downstream of each ORF is located a polyadenylation signal-encoding sequence which in this case is derived from the Simian virus 40 (SV40) early gene and is labelled SV40 polyA site in the diagram of Figure 1.

The term enhancer/promoter is used herein to denote combined or fused enhancer and promoter nucleotide sequences.

Following recombination, the above-mentioned mammalian cell enhancer/promoter drives expression of the *tk* gene. Possible alternatives include the promoter of the chicken B-actin gene, a tissue specific promoter such as for example the prostate specific antigen promoter and its associated enhancer region, or a tumour specific promoter such as for example the HSP-70

promoter, the MDR-1 promoter, the E2F-1 promoter or telomerase-related promoters.

As pointed out, other possible tumour sensitizing genes that may be used include the *E.coli* nitroreductase or cytosine deaminase genes, and the 5 mammalian cytochrome p450 2El or 2DVI cDNAs. Possible alternatives for the above-mentioned radiation-responsive promoter for the recombinase gene include promoters comprising different numbers of radiation-responsive elements, which may be the same or different, arranged in tandem, the entire enhancer/promoter DNA sequence, or only the enhancer DNA sequence of the 10 *egr-1* gene. Other alternatives include the sequences of the *TNF α* gene, the *Nfk β* gene, the *c-fos* gene, the *jun-b* gene, the *c-jun* and the *c-myc* gene or their functional equivalents. Where a heat-responsive promoter or expression control element is used, as previously mentioned this can be the promoter of the *hsp-90* gene or its functional equivalent.

15 In Figure 1 the region labelled "Neo gene" indicates the amino-glycoside phosphotransferase gene ORF complete with a mammalian cell promoter and SV40 polyadenylation site, which is incorporated for selective growth of transfected mammalian cells *in vitro* in the selective antibiotic G418. This allows the vector to be used as a model to demonstrate aspects of the 20 invention *in vitro* in cultured human cells since it allows permanent cell lines to be established. Such cell lines, e.g. the pCE cell line (see later), can be used not only to demonstrate the working of the invention *in vitro* but they can also be used to produce tumour xenografts in immune-deficient mice for demonstrating the working of the invention *in vivo*. Also as shown in Figure 1, the plasmid 25 contains an antibiotic resistance gene expression cassette such as that conferring resistance to ampicillin in *E.coli*, (labelled Amp in Figure 1), a bacterial origin of replication for growth of the plasmid in *E.coli* (labelled *E.coli* ori in Figure 1), and a mammalian cell origin of replication for

replication of the plasmid in mammalian cells (not shown). The latter would be ideally located outside the region excised by the recombinase so that additional copies of the inactive plasmid might be produced by DNA replication prior to recombination. and following recombination additional copies of the activated 5 plasmid, now expressing the tumour killing or sensitizing gene(s), might also be produced by DNA replication, resulting in increased expression of the tumour killing or sensitisation gene(s).

The vector illustrated in Figure 2 (pComplete2) contains the same elements as are shown in Figure 1 but the region between the *loxP* sites has 10 been modified to contain another copy of the recombinase Cre ORF and of the SV40 poly A site, the direction of transcription of these sequences being opposite to that of the *tk* gene. Another copy of the CMV I.E. enhancer/promoter is located adjacent to the Stop cassette and there is another 15 intron located transcriptionally upstream of this. These elements are positioned such that excision and recombination of the region between the *loxP* sites by Cre (supplied by a radiation-upregulated Cre expression cassette located elsewhere in the plasmid as shown in Figure 2) results in the production of two circular plasmid molecules, the *tk* gene being expressed from one, while the *cre* gene in the other is now able to be expressed as a result of the repositioning of 20 the CMV I.E. enhancer/promoter upstream of the Cre ORF. This should ensure that any copies of the switchable plasmid which are present in the cells but which have not become activated by the initial removal of the Stop cassette would then be activated, without the requirement for additional doses of radiation or diathermy. Again, the Cre-expressing circularised fragment can be 25 designed to contain a mammalian cell origin of replication so that additional copies of the activated plasmid might be produced by DNA replication, resulting in increased expression of Cre.

The vector in Figure 3 (pComplete3) is designed so that that excision of the Stop cassette region between the *loxP* recombination sites generates a DNA fragment in which two different tumour sensitizing (or killing) genes flanking an internal ribosome entry site (IRES) come under the influence of a mammalian cell enhancer/promoter or tumour-specific enhancer/promoter also contained within that fragment, giving rise to continuous expression of both these two genes. The example shown in Figure 3 is of the *tk* gene and the E.coli nitroreductase gene (labelled Nitroreductase-ORF) that would be expressed by virtue of the CMV I.E. enhancer/promoter, following Cre-mediated recombination.

It will be appreciated that many variations of the above-described vectors can be produced. For example, in pComplete2 the positions of the Cre ORF and *tk* ORF flanking one of the *loxP* sites can be reversed so that, following the action of Cre, the shorter region between the *loxP* sites will express the tumour sensitizing gene whilst the residual vector will continuously express Cre. Furthermore, additional IRES will allow the expression of more than one protein or other tumour cell killing molecule from both of the vector fragments following recombination and both fragments might contain a variety or mixture of the Cre ORF and many other tumour sensitizing genes, and/or tissue or tumour-specific promoters or enhancer/promoters.

Vector Construction

By way of example, the manner of construction of the pComplete1 vector in accordance with the invention will now be described in somewhat greater detail.

The elements of the vector are assembled in standard ways well-known in the art of recombinant DNA technology from components available commercially or that can be made synthetically or derived by PCR-amplification of human DNA, and appropriate restriction endonuclease sites are

introduced which can be used to produce fragments with compatible termini. As hereinafter more fully described, these may then be ligated together as required.

The components of the vector(s) in this specific example are
5 conveniently obtained as summarised below:

(a) The or each tandem array of radiation responsive promoter elements corresponding to the radiation responsive promoter element of the *egr-1* gene is conveniently synthesised initially as a short single-stranded oligonucleotide as described by Weichselbaum *et al.* (1994) "Gene therapy targetted by radiation preferentially radiosensitizes tumour 10 cells". *Cancer Research*, 54, 4266-4269, the content of this paper being incorporated herein by reference. The terminal regions of this single-stranded oligonucleotide can be designed so that, following annealing to a suitable complementary oligonucleotide, the resulting double-stranded molecule can be ligated into the plasmid vector pCI-*neo* (commercially available from Promega) following its digestion 15 with appropriate restriction endonuclease/s. in this case *Bg*III and *Sg*I.

(b) The Cre recombinase protein coding sequence of the control gene is readily obtained as a *Xho*I/*Mlu*I fragment from the commercially 20 available plasmid pBS185 (Gibco BRL™ - Life Technologies, Inc.).

(c) The cytomegalovirus (CMV) immediate-early (I.E.) enhancer/promoter, SV40 polyadenylation signal, the *E.coli* and mammalian cell origins of replication, ampicillin resistance gene and aminoglycoside resistance cassette (comprising the CMV I.E. 25 enhancer/promoter, aminoglycoside phosphotransferase ("neo") protein coding sequence and SV40 polyadenylation signal) are also conveniently obtained from the plasmid vector pCI-neo commercially available from Promega.

(d) The cytomegalovirus (CMV) immediate-early (I.E.) enhancer/promoter, and a suitable multiple cloning site (MCS) is obtainable as a *Bam*Hi/*Bgl*II fragment from plasmid pCI, also commercially available from Promega.

5 (e) The *loxP*-flanked Stop cassette can be obtained as an *Eco*Ri/*Spe*I fragment from GibcoTM plasmid pBS302.

(f) The *Herpes Simplex* virus *tk* gene has the nucleotide sequence originally published by Wagner, M.J. *et al.* (1981) "Nucleotide sequence of the thymidine kinase gene of herpes simplex virus type-1", *Proc. Natl. Acad. Sci. USA*, 78, 1441-1445 and for the purpose of the present invention it is conveniently cloned by standard methods into plasmid pBR322 (Gibco) to produce a plasmid vector here termed pBR-*tk* which provides a carrier and reservoir of the gene.

10 More specifically, the production of the vector pCompletel shown in Figure 1 from synthetic oligonucleotides, plasmids pCI-*neo* and pCI (Promega), plasmids pBS302 and pBS185 (Gibco) and the above-mentioned pBR-*tk* plasmid can be carried out in a series of stages substantially as described below with reference to the scheme illustrated in the diagram of FIGURE 4, and with reference to the oligonucleotide sequences shown in TABLE 1 at the end of the 15 present description.

15

20

In the diagram of FIGURE 4, the double-framed boxes indicate starting materials, and the various stages may be carried out substantially as follows:

(1) The CMV I.E. enhancer fragment is removed from plasmid pCI-*neo* following digestion with *Bgl*II and *Sgf*I and is replaced with a double stranded oligonucleotide labelled "RR-elements(a)". This contains a tandem array of 6 radiation-responsive elements and is produced by

25

annealing synthetic oligonucleotides EGRE1 and EGRE2 (see Table 1) to generate the plasmid, labelled pEGRL(a)-neo.

(2) The *cre* recombinase gene protein coding region, obtained as a *Xho*I/*Mlu*I fragment of pBS185, is inserted into *Xho*I/*Mlu*I-digested pEGRL(a)-neo to generate pEGRL(a)-*cre*.

(3) The *Not*I site in the pEGRL(a)-*cre* Multiple Cloning Site (MCS) is deleted by *Not*I digestion, then the resultant single-stranded termini filled-in using the Klenow enzyme, followed by self-ligation of the plasmid.

10 (4) The CMV I.E. promoter/enhancer along with its associated multiple cloning site is obtained as a *Bam*HI/*Bgl*II digest fragment from plasmid pCI, and is ligated into the *Not*I-deleted pEGRL(a)-*cre*, following partial digestion of the latter with *Bam*HI, thereby to produce pEGRL*cre*MCS2.

15 (5) The Stop cassette, obtained from the Gibco™ plasmid pBS302 by digestion with *Eco*RI and *Sph*I, is ligated into *Eco*RI/*Xba*I digested pCI-*neo* plasmid to generate pStop.

(6) The *tk* gene protein coding region is obtained as a PCR amplification product of pBR-*tk* using the primers *Clatk* (SEQ ID No: 9) and *tkNot* (SEQ ID No: 10) shown in TABLE 2 at the end of the present description. Following *Cla*I and *Not*I digestion the PCR product is ligated into *Acc*I/*Not*I-digested pStop to generate pStop-*tk*.

20 (7) The *Not*I/*Eco*RI fragment of pStop-*tk* containing the Stop cassette and the *tk* ORF is ligated into *Eco*RI/*Not*I-digested pEGRL*cre*MCS2 to generate pComplete1.

Radiation Switched Operation

Figure 5 illustrates in relation to pComplete 1 the molecular switching process in which Cre recombinase transcription and translation is induced following exposure to ionising radiation. The Cre enzyme expressed acts upon 5 pComplete1 to excise the Stop cassette by recombination at the *loxP* sites. This results in the formation of a small circular DNA fragment containing the Stop cassette and a *loxP* site and of a recircularised plasmid (pCompletel: activated form) that transcribes and translates the thymidine kinase DNA sequence. The expression of this enzyme can then in turn cause activation of the prodrug. 10 gancyclovir (provided by systemic injection) which will kill host cells and also cells in contact with the host cells (via the bystander effect).

Modified Vectors

Alternative vectors containing different promoter and/or enhancer elements can be constructed by analogous methods. Thus, as indicated earlier, 15 the vectors may be constructed so as to contain the entire radiation responsive *egr-1* enhancer/promoter. This comprises nucleotides -676 to +10 according to Sakamoto *et al.* ("5' Upstream sequence and genomic structure of the human primary response gene *egr-1/TIS8*" (1991) *Oncogene* 6, 867-871) and can be isolated from human DNA by PCR amplification using the primers EGRE6 20 (SEQ ID No: 7) and EGRE5 (SEQ. ID NO: 6) shown in TABLE 2. The enhancer region alone (nucleotides -676 to -178) can be isolated from human DNA by PCR amplification using the primers EGRE5 (SEQ. ID NO: 6) and EGRE7 (SEQ. ID NO: 8) shown in TABLE 2. These primers, which are designed to introduce appropriate restriction endonuclease or other suitable 25 sites for cloning, are conveniently synthesised using commercially available oligonucleotide synthesising apparatus. The human DNA required for carrying out the PCR amplification of these radiation responsive sequences is conveniently isolated from HeLa cells by a conventional method, as described

for example by Sambrook, J., Fritsch, E. F. & Maniatis, T. (1989), "Molecular Cloning - A Laboratory Manual", *Cold Spring Harbor Laboratory Press, U.S.A.*).

Alternative recombinase control genes and target sites as well as other
5 tumour sensitizing mechanisms can also be used. As previously mentioned, one example of the latter is the *E.coli* nitroreductase gene in respect of which the protein coding sequence (nucleotides 166 to 831) can be isolated by PCR amplification of *E.coli* strain B DNA using oligonucleotide primers designed to introduce suitable restriction endonuclease sites for cloning purposes.

10 In modifications in which the components of the vectors are arranged in different ways in order to achieve the required trigger and switching effect of radiation (or other expression inducing influences), as has been described in relation to the vector illustrated in the diagram of Figure 2 the region between the recombinase (*loxP*) target sites can contain a duplicate copy of the
15 recombinase gene and another mammalian cell promoter on opposite sides of the Stop cassette, the arrangement being such that the recombination and re-orientation brought about by the action of the recombinase initially expressed will bring this second copy of the recombinase gene under the influence of this other promoter, thereby resulting in the continuous expression of this second
20 copy of the recombinase coding sequence. As has already been pointed out, continued expression of the recombinase enzyme after triggering by a single dose of radiation will ensure that further radiation or heat treatment will not be required if there were to be inadequate or inefficient recombination of the vectors in the tumour cells due to low levels of heat or radiation-induced
25 expression of the recombinase after the initiating dose of radiation or heat. In this way the dose of heat or radiation or other expression inducing influence required to trigger the removal of the Stop cassette and the activation of the silenced expression cassette or tumour sensitizing gene(s) in every copy of the

vector taken up by the cells can be kept minimal. This can be particularly important if the system of vector delivery results in the uptake of more than one copy of the vector into the host cells or if the vector undergoes replication before recombinase-mediated recombination.

5 Additionally, multiple killing or sensitisation factors can be expressed in a similar way by means of internal ribosome entry sites (IRES) that allow the expression of more than one protein coding sequence under the influence of a single promoter region, as has been described in connection with the vector of Figure 3. This can allow, for example, the simultaneous expression of different prodrug activating proteins and immune response stimulating factors, or different combinations thereof, or different combinations of prodrug activating proteins and other proteins, including Cre, or other molecules that kill cancer cells. It is also possible to express fusion proteins having more than one functional prodrug activating activity.

10 The vectors may be designed to be episomal, requiring for example an Epstein-Barr virus nuclear antigen-1 gene element and origin of replication, or a SV40 origin of replication. Alternatively, they may be designed to integrate into host cell DNA as would be the case for retroviral or adenoviral vectors. The excised region between the recombinase target sites or the region outside 15 the recombinase target sites encompassing the tumour killing or sensitizing gene(s) may also be designed to be episomal and to have its own mammalian cell origin of replication. It is recognised in some cases that the capacity of certain viral genomes to harbour DNA is size-limited and therefore some components of the vectors herein illustrated may need to be eliminated in order 20 that the essential genetic information can be accommodated.

Testing

The operation of the system can be tested in cultured human tumour cells and xenografts established therefrom using vectors of the general kind herein described.

5 (a) Demonstration of radiation-mediated switching in human tumour cells

In order first to prove the "molecular switching" principle of the invention, a series of experiments was carried out to test the operation of the various components individually and then in combination. For convenience and for the purposes of accurate quantitaton of any effect observed. instead of a 10 prodrug activating gene, the gene encoding the Green Fluorescent Protein (GFP) from the jellyfish *Aequorea victoria* was used as a sensitizing "reporter" gene. The ORF of this gene was cloned into the vector pStop (see Figure 6) downstream of the Stop cassette to generate pStop-gfp. This plasmid, which should not be able to express GFP unless it undergoes recombination, was 15 transfected into MCF-7 cells in order to provide a control.

In constructing pStop-gfp, the Green Fluorescent Protein (GFP) open-reading frame was excised from plasmid pEGFP-1 (obtained from Clontech) with *Cla*I and *Not*I and was cloned into the *Acc*I/*Not*I sites of pStop. The radiation-responsive synthetic *Egr-1* enhancer (E4) formed by annealing 20 EGRE3 and EGRE4 (see Table 1) containing four repeats of the previously defined decanucleotide radiation-responsive expression control elements (see Datta *et al.*, (1992) *Proc. Natl. Acad. Sci. U.S.A.* 89, 101149-101153) was cloned into the pCI-neo vector to generate pEGRL(b)-neo. The *cre* gene was obtained from pBS185 as an *Xho*I/*Mlu*I fragment and was ligated into 25 *Xho*I/*Mlu*I-digested pEGRL(b)-neo to generate pEGRL(b)-cre. This, together with the plasmid pStop-gfp, was transfected into MCF-7 cell lines using the cationic lipid transfection reagent Lipfectamine. pStop-gfp-transfected and pStop-gfp plus pEGRL(b)-cre-transfected cells were irradiated (5Gy and 10Gy

in two 5Gy doses), and the number of fluorescent cells within a constant aliquot of the transfected population was measured by FACS, as hereinafter described. As can be seen in Figure 7, transfection with the pStop-*gfp* vector alone resulted in very low numbers of fluorescent cells, although there was a slight
5 increase in this number following irradiation. Transfection with pStop-*gfp* plus pEGRL(b)-*cre* resulted in a larger number of fluorescent cells than with pStop-*gfp*-transfected irradiated cells, but this number was further increased to a much greater extent following irradiation. It can thus be concluded that irradiation activated the radiation-responsive promoter in pEGRL(b)-*cre* which resulted in
10 the expression of Cre recombinase which in turn recombined the pStop-*gfp* vector, causing expression of GFP. This therefore demonstrated the operation of the general "switch" principle described in connection with this invention.

(b) Demonstration of radiation-mediated killing in human tumour cells

To test this aspect, MCF-7 cells were transfected as described above
15 with the plasmid pEGRL(b)-*cre* and a permanent cell line, pCE, was established by G418 selection. This was transfected with the plasmid pStop-*tk* described earlier and the cells were plated in 96-well microtitre plates in either complete tissue culture medium (RPMI + 10% foetal calf serum) or the same containing 50 mM ganciclovir (GCV). Eight hours after plating the cells were
20 irradiated as an attached monolayer (^{60}Co gamma rays, approx 1Gy/min) at 37°C. MTT assays were performed 24 hours later and radiation survival data were presented as a mean (\pm SD) of 48 replicates per dose point.

As shown in Figure 8 there was some degree of cell killing with increasing doses of radiation in the absence of GCV. However, in the presence
25 of GCV, there was substantially more killing at doses of 2Gy or more. This is consistent with radiation-mediated upregulation of the *cre* recombinase gene in the pCE plasmid and subsequent recombination of the *loxP* sites, resulting in the removal of the stop cassette, leading to the synthesis of thymidine kinase

and subsequent action of the latter on GCV to convert it to a cytotoxic metabolite. The absence of an effect at the low dose of 1Gy is also consistent with this being below a threshold dose for activation of the radiation-responsive promoter in the *cre*-encoding vector. The increased cell killing at 5Gy may
5 reflect synergism between radiation and activated GCV.

(c) Further testing and demonstration of radiation-mediated gene therapy treatment of tumour cells

In another series of experiments, a cell line MCF-7/E4*cre* was established that constitutively contained a radiation-inducible *Cre* recombinase gene. MCF-7 cells were transfected with a plasmid designated pE4*cre* containing a *Cre* gene and the synthetic *Egr-1* enhancer/promoter E4. These cells were then subjected to G418 selection. Cell lines were cloned by serial dilution in G418 (0.5 mg mL⁻¹). The same technique was used to establish a cell line designated MCF-7/E4*tk* composed of MCF-7 cells transfected with a *tk* gene containing plasmid pE4*tk*. MCF-7 cells and cell lines were transfected with pE4*cre*, p*Stk* or p*tk* plasmids, p*Stk* being a plasmid containing a Stop cassette interposed between a *tk* gene and the strong constitutive cytomegalovirus immediate early gene promoter (CMV IE). Plasmid p*tk* was similar, but without the Stop cassette, i.e. a plasmid containing a *tk* gene under
15 the direct control of a cytomegalovirus gene CMV IE promoter.
20

After transfection, cultures were irradiated 3-4hrs later, and after 5 days cell growth was determined, calculated in the presence and absence of 50 µM GCV (ganciclovir) for all cell lines in independent experiments.

The results illustrated in Figure 9a and 9b show that the molecular
25 switch is very efficient. Using the MCF-7/E4*cre* cell line as a control, (O in Figure 9a), similar increases in the extent of GCV-mediated growth inhibition were seen following irradiation of MCF-7 cells that had been transfected with the plasmid p*tk* in which the CMV promoter operated the expression of tk,

(graph ▼ in Figure 9a) and MCF-7-*cre* cells that had been transfected with pStk wherein the radiation-triggered molecular switch was required to achieve GCV-mediated sensitisation (graph ● in Figure 9a). In comparison, using MCF-7 cells as a control MCF-7 cells transfected with pE4tk (in which the synthetic radiation-responsive promoter E4 directly operated the expression of a *tk* gene) the increase in sensitisation following irradiation was considerably less (graph ■ in Figure 9b).

Since the switch system resulted in the same levels of sensitisation as an unswitched system, it is concluded that the amounts of Cre recombinase produced upon irradiation, even at the lowest dose used, were not a limiting factor in the effect. For clinical treatment of tumours, it would thus be an advantage to administer mixtures of small amounts of Cre-expressing plasmid together with larger amounts of tumour-sensitizing gene expressing plasmid or plasmids in order to obtain the maximum molecular switching effect and hence 15 tumour sensitisation from a single dose of radiation.

It will be appreciated that the Stop cassette-containing plasmids may themselves be a mixture containing different tumour sensitizing genes and in practice a mixture most suitable for the tumour type being treated would be used. This would facilitate the manufacture of the plasmids since the cre-expressing plasmid would be common to any treatment protocol and would be combined with a stop cassette plasmid of various kinds such as would be most appropriate for the specific characteristics of the tumour in question.

Demonstration of the use of Platinum containing drugs as control gene
25 expression inducing agents

As indicated earlier, it has also been found that genes having radiation responsive promoters or other expression regulatory elements can in many

cases respond also to, and be upregulated by, platinum containing drugs such as cisplatin (CDDP) for example. This is demonstrated in Figure 10 which shows the results obtained in experiments in which the expression was determined (by FACS analysis) of Green Fluorescent Protein (GFP) encoded in a plasmid vector pE4-*gfp* containing the *gfp* gene directly controlled by the synthetic radiation responsive promoter E4.

As shown in Figure 10, 1micromolar CDDP induced the expression of GFP to a similar extent as did ionising radiation (IR). Thus CDDP was able to induce gene expression regulated by the radiation-responsive elements in the promoter region upstream of the *gfp* gene. Hence this agent or, in principle, any similar agent or agents producing similar or effective DNA damage could be used to trigger the molecular switch and effect the expression of tumour sensitizing genes. It was noted that combinations of CDDP and IR were more effective than either of the treatments alone. Thus two or more agents may synergise and allow the switch effect to occur at even lower sub-toxic and sub-lethal doses of the individual inducing agents.

To use such platinum containing drugs, or indeed any chemical inducing agents of which there are many possibilities (given an appropriate choice of control gene promoter elements), to provide the required control gene expression inducing influence, it is anticipated that for clinical administration such agents would most probably be encapsulated into cationic liposomes with a view to tumour targeting. Thus, such targeted drugs may be used instead of radiation in the present invention. Importantly, and by analogy with the radiation-mediated gene therapy, the amounts of these chemical inducing agents that would be required to trigger the response would generally be less than those required to kill the tumour cells if the agents were to be used without the molecular switch.

Demonstration of Control gene expression induced by Hypoxia

In these experiments the HIF-1 binding domain of the enolase-1 gene was produced as a double-stranded oligonucleotide by annealing of the synthetic oligonucleotides HRE1 and HRE2 (see Table 3). The annealed 5 oligonucleotide was ligated into *Bgl* II-*Sgf* I digested pCIneo (Promega) to produce a plasmid designated pHCIneo. pHCIneo was digested with *Eco*RI and *Not*I and the *Eco*RI-*Not*I fragment of pEGF-1 (Clontec) that contains the GFP gene was ligated into the resulting gap to generate a vector pHRE-EGFP.

NB1G human neuroblastoma cells were plated in culture medium at 10 7.5 x10⁴ cells per well in a 24-well plate and incubated at 37°C in an atmosphere containing 20% oxygen. Three days later they were transfected with pEGFP (which contains no promoter), pCMVGFP (which contains the CMV promoter) or pHREGFP (which contains the hypoxia-responsive promoter) using as a transfecting agent Lipofectamine Plus as described by the 15 manufacturer. After 24 hours, the medium was replaced with fresh medium or medium containing 25 micromolar CoCl₂. Following incubation overnight in an atmosphere containing 5% O₂. FACS analysis was performed as described elsewhere. These conditions were intended to produce hypoxia in the transfected neuroblastoma cells.

20 The results are illustrated in Figure 11 which shows that pHREGFP-transfected cells were more fluorescent than the control pEGFP or pCMVGFP cells under low (5%) oxygen conditions, whilst CoCl₂ had no or only a slight inhibitory effect. However, in the latter two cases, with the pHREGFP-transfected cells there was a considerable increase in the number of fluorescent 25 cells in the population. Thus it was shown that GFP expression from the pHREGFP vector can be induced by conditions of hypoxia.

METHODS and MATERIALS - Summary

For completeness there now follows an outline or summary of some of the methods, techniques and materials which have generally been used in development and testing of this invention unless stated otherwise.

Cells

5 All bacterial cloning was carried out in *E.coli* strain XL-1 blue MRF' (Stratagene).

The human cell line used for transfection was MCF-7 (breast carcinoma) obtained from the American Tissue Culture Collection (HTB No. 22). The DNA used for PCR amplification of the *egrf-1* promoter elements was
10 extracted from the human cervical carcinoma line, HeLa.

Plasmids

The pCI and pCI-neo plasmids were obtained from Promega.

The pEGFP-1 plasmid was obtained from Clontech.

15 The pBS185 (containing the *cre* recombinase gene) and pBS302 (Stop cassette) plasmids were obtained from Gibco BRL™ (Life Technologies, Inc.).

The plasmid clone pBR-*tk* was produced by cloning the nucleotide sequence of the *Herpes simplex* virus type 1 (HSV) thymidine kinase (*tk*) gene, strain CL101. (described by Wagner *et al.*, (1981) as previously mentioned) into Gibco plasmid pBR322, using standard methods.

20 Enzymes

All restriction endonucleases were obtained from MBI Fermentas, with the following exceptions. *SgfI* and I-*Ppo*-I were from Promega. *AccI*, *Ascl* and *SpeI* were from New England Biolabs.

25 T4 DNA ligase, Taq DNA polymerase and Klenow were from MBI Fermentas. Expand High Fidelity polymerase was from Boehringer Mannheim. Taq Plus Precision was from Stratagene.

All enzymes were used in the buffers supplied and in accordance with the manufacturers instructions.

DNA Purification

High molecular weight HeLa DNA was extracted from cultured cells 5 using the MBI Fermentas Genomic G2 kit. Large-scale plasmid DNA preparations were carried out using the Qiagen Maxi-purification kit.

DNA extraction and purification from agarose gels (Seakem GTG agarose, FMC Bioproducts) was carried out using the Qiagen™ Gel Extraction kit.

10 After restriction endonuclease digestion or modification, DNA was routinely purified using the Nucleon PCR/Oligo Clean-up kit.

Nucleotides

Deoxynucleotides for PCR were obtained from Pharmacia and used at 15 0.2mM final concentration.

Synthetic oligonucleotides for providing the radiation responsive elements and PCR primers less than 60 base pairs in length were obtained from Gibco BRL™. Those of a greater length were obtained from Cruachem.

Transformation of *E.coli*

20 The procedure used to prepare competent cells for molecular transformation and freezing was that outlined in the previously mentioned reference of Sambrook *et al.* (1989), with a 10mL 0.1M magnesium chloride cell pellet resuspension step prior to the first calcium chloride stage to increase transformation efficiency. The transformation procedure itself was also from 25 this source.

Growth and transfection of human cells

Pre-confluent human tumour cells (MCF-7 [ATCC HTB 22]) grown in RPMI tissue culture medium + 10% foetal calf serum (FCS) were washed with PBS, serum free media (SFM) and then exposed to a transfection mixture of 15 μ L lipofectamine (Gibco BRLTM) mixed with 85 μ L double-distilled water 5 (ddH₂O) combined with 10 μ L (5 μ g) of plasmid DNA mixed with 90 μ L ddH₂O. This addition was immediately followed by adding 800 μ L RPMI + 5% FCS and incubating at room temperature for 5 hours. Subsequently, 1mL of RPMI + 15% FCS was added for 21 hours, after which the medium was removed and replaced with fresh complete RPMI + 10% FCS.

10 Irradiation

Cells were irradiated at 37°C with 5Gy and 10Gy (2 x 5Gy) of cobalt-60 (gamma-rays) at a dose rate of 1 Gy min⁻¹.

FACS analysis

Green Fluorescent Protein (GFP) expression was assessed by 15 fluorescent activated cell sorting (FACS) and scanning (Becton-Dickinson FACScanTM; excitation 488 nm). Monolayers of cells were trypsinised, washed with PBS (phosphate buffered saline) and finally resuspended in PBS as a single cell suspension. FITC fluorescence was measured at 530 \pm 15 nm, and cell debris resulting in forward and side scatter was gated out. The gated 20 fluorescence 1 profile of the FACScanTM software was used to determine the number of fluorescent cells in the sample.

MTT assay

Cytotoxicity assays were performed using a modification of a method previously published (Morten *et al.*, 1992 "Upregulation of O⁶-alkylguanine-DNA-alkyltransferase expression and the presence of double minute chromosomes in alkylating agent selected Chinese hamster cells" 25 *Carcinogenesis* 13(3): 483-487). Briefly, confluent cultures of cells were

trypsinised, twice washed in PBS and resuspended to 2000 cell/mL in complete tissue culture medium (RPMI + 10% foetal calf serum) containing 50 mM ganciclovir (GCV). Aliquots (200 µL) of this suspension were pipetted into each well of 96 well microtitre plates. The plates were incubated at 37°C for 4 days, after which the medium was aspirated and replaced with 100 µL of a 3 mg/mL solution of MTT [3-(4,5-dimethyl-thiazol-2-yl)-2,5-diphenyltetrazolium bromide] in PBS for 3 hours, followed by 200 µL of DMSO. The plates were agitated to ensure complete dissolution, and were read on a multiplate reader (Flow Ltd.) at 530 nm and 690 nm.

10 DNA Manipulations

(a) *PCR:*

The *tk* ORF was obtained by PCR using the specific primers shown in TABLE 2. The proof-reading DNA polymerases, Expand High-fidelity and Taq Plus PCR were used to avoid introduction of mutations. After 2 minutes of initial DNA strand melting at 94°C, polymerase was added. Amplification was carried out for 15 cycles under the following conditions: 94°C 1 min, 55°C 1 min, 72°C 1min. The latter, elongation, step was extended by 20 seconds each cycle for the last 5 cycles. A single 5 min elongation was also added as a final step. Products amplified from plasmid DNA were agarose gel purified to remove the original template.

20 (b) *DNA Restriction endonuclease digestion:*

As already described, a series of steps was needed to produce the pComplete1 construct. Briefly, to recap. and summarise:

25 (i) The Stop cassette was excised from pBS302 with *Eco*RI and *Spe*I and was cloned into the *Eco*RI and *Xba*I sites of pCI-*neo* to produce pStop (see Figure 4).

(ii) The *tk* ORF was amplified from the HSV plasmid clone pBR-*tk*, using the *tk* PCR primer pair shown in TABLE 2. The purified PCR fragment was then treated with *Cla*I and *Not*I and cloned into *Acc*I/*Not*I digested pCI-*neo* to generate pStop-*tk* (also see Figure 4).

(iii) The radiation-responsive *egr-1* elements (RR-elements(a)) were cloned into the pCI-*neo* plasmid as described, using the *Bgl*II and *Cgt*I sites to generate pEGRL(a)-*neo*.

(iv) The Cre recombinase ORF was excised from pBS185 with *Xba*I and *Mlu*I and cloned into those sites in pEGRL(a)-*neo* to generate pEGRL(a)-*cre*.

(v) The *Not*I site was removed from the pEGRL(a)-*cre* plasmid by *Not*I digestion. This was followed by "filling-in" of single-stranded overhangs using Klenow, then re-ligation.

(vi) The *Bgl*II/*Bam*HI fragment of the pCI plasmid, containing the CMV I.E. enhancer/promoter, multiple cloning site and SV40 polyadenylation site, was isolated and then cloned into the *Not*I-deletion construct via partial *Bam*HI digestion, allowing insertion downstream of the neomycin resistance gene to generate pEGRL-*creMCS2*.

(vii) Finally, the *Eco*RI/*Not*I Stop-*tk* cassette of pStop-*tk* was cloned into the inserted multiple cloning site in pEGRL-*creMCS2* to produce pComplete1.

(c) *Ligation:*

For ligations involving only single-stranded terminal overhangs, 20-50ng of plasmid vector DNA was added to 60-200ng of potential insert DNA

and incubated at 21-25°C for approximately 20 hours before transformation. Blunt-end ligations were performed at 4°C.

The radiation responsive element containing oligonucleotides were annealed by placing 0.2-0.5nmoles of each complementary molecule in a 5µL total volume and heating to 55°C for 5min before leaving to cool for about 24hrs. This mixture was then added to 50-100ng of plasmid vector for ligating as normal.

CLINICAL APPLICABILITY

In clinical use the vector material of the present invention will be administered to a cancer patient in a suitable dose and in a suitable pharmaceutical composition using, as already explained, either a virus-based or virus-free method of DNA delivery.

After administration of the composition containing the vector to the patient an appropriate time is allowed for uptake and incorporation into cells. The patient's tumour may then be subjected to a suitable dose of diathermy, ionizing radiation or other appropriate exogenous inducing agent (unless relying on an endogenous inducing influence). In the case of ionizing radiation, this is preferably applied using conformal radiotherapy apparatus, or a tumour-targeting radiolabelled agent such as a tumour specific radio labelled antibody or cytokine may be used. In the case that the tumour has metastasised, when appropriate whole body diathermy or radiation or a tumour-targeting radiolabelled agent may be used. The inducing agent activates the inducing agent responsive promoter of the control gene with the result that recombinase protein is expressed. Although this expression may be transient, the action of the recombinase results in the recombination of the recombinase target sites and hence excision of the Stop cassette, resulting in the permanent expression of the tumour-sensitizing gene or genes as hereinbefore described in connection with the specific embodiments referred to.

When a prodrug activating enzyme is encoded by the sensitizing gene(s), usually after administration of the vector composition in the course of clinical use and application of heat or ionizing radiation or other inducing agent as the case may be, an appropriate therapeutically effective dose of the prodrug 5 is administered in a conventional formulation by any suitable route. This then brings about the death of the cells that express this prodrug activating enzyme, and also cells in the vicinity by virtue of the bystander effect.

In an alternative, if the tumour killing or sensitizing gene encodes proteins such as for example interleukin-2 (IL-2) or granulocyte-macrophage 10 colony-stimulating factor, an immune response will be stimulated in the host that should eradicate the tumour cells. On the other hand, if the tumour killing or sensitizing gene encodes a cytotoxic protein, e.g. ricin, expression can result directly in the killing of the host tumour cells.

In the case that the tumour sensitizing gene encodes a ribozyme or an 15 antisense RNA molecule, these would be designed to bind to and cleave, or elicit the cleavage of, specific messenger RNA molecules that encode proteins the non-production of which would lead to the death of the cells or render the cells more sensitive to killing by exogenous agents or ionising radiation. However, with this approach it is likely that only those cells to which the 20 therapeutic DNA is delivered would be killed or sensitised, whereas with prodrug activation bystander effects can occur.

The advantages of the invention include the fact that in some 25 embodiments using diathermic heat or ionizing radiation as exogenous inducing agents recombination and subsequent activation of the vector material only occurs in the region of diathermy or in the irradiated area. The uptake of the vectors into normal cells at distant sites will not therefore result in the ability of such cells to activate prodrugs or otherwise elicit tumour sensitisation. Furthermore, very low sub-therapeutic doses of ionizing radiation can be

effective in inducing expression of, for example, the *egr-1* enhancer/promoter. Hence, in the case that the tumour has metastasised, tumour-seeking radiolabelled agents that are inefficient in killing tumour cells *per se* can be used to instigate the recombination events and enable the activation of 5 prodrugs, or the other cancer cell-killing processes described, in potentially all metastatic sites as well as in the primary tumour.

As will be seen, the invention presents a number of different aspects and it should be understood that it embraces within its scope all novel and 10 inventive features and aspects herein disclosed, either explicitly or implicitly and either singly or in combination with one another. Also, many detail modifications are possible and, in particular, the scope of the invention is not to be construed as being limited by the illustrative example(s) or by the terms and expressions used herein merely in a descriptive or explanatory sense.

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TABLE 1

Synthetic single-stranded oligonucleotides used to produce double-stranded molecules containing radiation responsive elements (RRelements) E provided by repeats of the decamer: 5' CCTTATTG (SEQ ID NO:1).

5 Plasmid pComplete1 contains a total of 6 elements (E6) arranged in series in a tandem array:

EGRE 1

BgII(part of)

5' GATCTCCTTA TTTGGCCTTA TTTGGCCTTA TTTGGCCTTA
10 TTTGGCCTTA TTTGGCCTTA TTGGGCGAT

Sgfl(part of)
(SEQ. ID NO: 2)

EGRE 2 (complementary to EGRE1)

5' CGCCCAAATA AGGCCAAATA AGGCCAAATA
15 AGGCCAAATA AGGCCAAATA AGGCCAAATA AGGA
(SEQ. ID NO:3)

Examples of similar oligonucleotides containing four repeats (E4) of SEQ. ID NO: 1

20 EGRE 3

BgIII (part of)

5' GATCTTATT TGGCCTTATT TGGCCTTATT TGGCCTTATT
 TGGGCGAT

25 (SEQ. ID NO: 4)

EGRE 4 (complementary to EGRE 5)

5' CGCCCAAATA AGGCCAAATA AGGCCAAATA
AGGCCAAATA AGGA

30 (SEQ. ID NO: 5)

TABLE 2

A. Primers used for PCR amplification of promoter/enhancer sequences of human primary response gene *egr-1/TIS8*.

For enhancer/promoter:5 EGRE 5*Bgl*II

5' TCCAGATCTC CCGGTTCGCT CTCACGGTCC CTGAGG

(SEQ. ID NO: 6)

EGRP 610 *Asc*I

5' CGGCGCGCCG CTGGATCTCT CGCGACTCCC CG

(SEQ. ID NO: 7)

For enhancer alone:EGRE 715 *Sgf*I

5' ACTGCGATCG CGGGCCCGGC CC GGCCCCGCA TCCCAGGCC

CC

(SEQ. ID NO: 8)

B. Primers used for PCR amplification of Thymidine kinase gene

20 Clatk:*Cla*I

5' CCATCGATAT GGCTTCGTAC CCCGGC

(SEQ. ID NO: 9)

*tk*Not:25 *Not*I

5' AAGGAAAAAA GC GGCCGCCT CCTTCCGTGT TTCAGTTAGC

(SEQ. ID NO: 10)

TABLE 3

Synthetic single-stranded oligonucleotides used to produce double-stranded molecules containing the hypoxia responsive region of the Enolase-1
5 gene promoter.

HRE1

Bgl II (part of)

5' GATCTAGGGC CGGACGTGGG GCCCCGTAGG CACGCTGAGT
10 GCGTGCGGGGA CTCGGAGTAC GTGACGGAGC CCCGCGATGC
 GAT

Sgf I (part of)

(SEQ. ID NO:11)

15

HRE2 (complementary to HRE1)

5' CGCATCGCGG GGCTCCGTCA CGTACTCCGA GTCCCGCACG
 CACTCAGCGT GCCTACGGGG CCCCACGTCC GGCCCTA

(SEQ. ID NO:12)

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CLAIMS

1. Vector material characterised in that said material contains:
 - (a) a tumour cell sensitizing gene or genes of which expression in a tumour cell yields a sensitizing gene expression product having a potential to cause tumour cells to be killed and destroyed, or to be eliminated, or otherwise to be inactivated, or to be rendered sensitive and/or vulnerable to destruction;
 - (b) a sensitizing gene expression regulatory system, including promoter means, for said sensitizing gene or genes;
 - (c) at least one control gene; and
 - (d) a control gene expression regulatory system responsive in use in a transfected cell to the effect of a predetermined exogenous or endogenous expression inducing influence so as to induce expression of said control gene to yield an expression product having a capacity to establish an operative linkage between said sensitizing gene expression regulatory system and said sensitizing gene or genes effective to trigger and switch on or permit continuous or permanent expression of the latter to bring about continuous production of said sensitizing gene expression product.
- 20 2. Vector material as claimed in Claim 1 wherein the tumour sensitizing gene expression product is an enzyme or other bioactive agent that can convert a predetermined inactive prodrug into an active cytotoxic drug.
- 25 3. Vector material as claimed in Claim 2 wherein the tumour-sensitizing expression product is a prodrug activating enzyme selected from the following:
 - HSV thymidine kinase, nitroreductase, cytosine deaminase, cytochrome p450 2E1 or cytochrome p450 2DVI.

4. Vector material as claimed in Claim 1 wherein the tumour-sensitizing expression product is an immune-response modifying agent selected from the expression products of the following:

5 GM-CSF, IFN-alpha, IFN-beta, IFN-gamma, IL-1beta, IL-2, IL-4, IL-6, IL-8, IL-10, IL-12, IL-15, TNFalpha.

5. Vector material as claimed in any one of the preceding claims wherein the control gene encodes a recombinase enzyme that acts on recombinase target sites to modify the vector material to establish said operative linkage between the sensitizing gene expression regulatory system and the sensitizing gene or 10 genes.

6. Vector material as claimed in Claim 5 wherein the control gene and the recombinase target sites are part of a Cre-loxP or a Flp-frt site specific recombination system.

7. Vector material as claimed in Claim 5 or 6 wherein said recombinase 15 target sites are separated by a region containing a "stop" sequence of nucleotides that blocks or otherwise prevents expression of the sensitizing gene or genes until removed by the action of said recombinase enzyme.

8. Vector material as claimed in Claim 5 or 6 wherein the protein coding 20 regions of the sensitizing gene or genes are operationally separated from the said promoters and wherein said recombinase target sites are arranged such that recombination brings about the juxtapositioning of the sensitizing gene promoters and protein coding regions of the sensitizing gene or genes resulting in their expression.

9. Vector material as claimed in any one of Claims 5 to 8 wherein the 25 control gene is a fusion gene that when expressed produces a fusion protein consisting of a recombinase and an intercellular trafficking protein (such as for example the virion protein VP22).

10. Vector material as claimed in any one of Claims 5 to 9 wherein the region between said recombinase target sites contains a duplicate copy of the recombinase control gene together with an associated promoter.
11. Vector material as claimed in any one of the preceding claims wherein 5 the sensitizing gene expression regulatory system incorporates at least one expression inducible element responsive to the effect of a predetermined exogenous or endogenous expression inducing influence.
12. Vector material as claimed in any one of the preceding claims wherein 10 the sensitizing gene is a fusion gene that when expressed produces a fusion protein consisting of a sensitizing protein and an intercellular trafficking protein.
13. Vector material as claimed in any one of Claims 1 to 8 wherein the or 15 each tumour sensitizing gene is selected from the *E.coli* nitroreductase gene, cytosine deaminase (CD) gene, *Herpes simplex* virus thymidine kinase (HSV-*tk*), mammalian cytochrome p450 2E1 or 2DVI gene, and their functional equivalents.
14. Vector material as claimed in any one of the preceding claims wherein 20 the tumour cell sensitizing gene or genes and the control gene are in separate vectors.
15. Vector material as claimed in any one of Claims 1 to 13 wherein the 25 tumour cell sensitizing gene or genes and the control gene are in the same vector.
16. Vector material as claimed in any one of the preceding claims for use in antitumour therapy characterised in that the use comprises the introduction of 25 the material into tumour cells.
17. Vector material as claimed in Claim 16 wherein said expression inducing influence is endogenous and tumour related, being produced by

tumour cells associated specifically with tumours to which said antitumour therapy is directed.

18. Vector material as claimed in Claim 17 further characterised in that at least one element of the control gene expression regulatory system is selected so that the control gene is automatically upregulated to an effective operational level when the vector material is introduced into cells of said tumours.

19. Vector material as claimed in Claim 18 wherein the selection of said at least one element of the control gene expression regulatory system is carried out using gene array technology.

20. Vector material as claimed in any one of Claims 17 to 19 wherein said control gene expression regulatory system responds in use in a transfected cell to said expression inducing influence where the latter is provided by a change in environmental thermal conditions in cells containing the vector material.

21. Vector material as claimed in any one of Claims 17 to 19 wherein said control gene expression regulatory system responds in use in a transfected cell to said expression inducing influence where the latter is provided by a change in local oxygen concentration.

22. Vector material as claimed in any one of Claims 16 to 19 wherein said control gene expression regulatory system includes an expression control element responsive in use in a transfected cell to a hypoxia condition in the cellular environment.

23. Vector material as claimed in Claim 16 wherein said control gene expression regulatory system responds in use in a transfected cell to said expression inducing influence where the latter is provided by an exogenous expression inducing agent applied to cells into which the vector material is introduced.

24. Vector material as claimed in Claim 23 wherein the dose level of said exogenous expression inducing agent needed to trigger and switch on expression of said sensitizing gene or genes in the cells of tissue into which the vector material has been introduced is substantially sub-lethal.

5 25. Vector material as claimed in Claim 23 or 24 wherein said control gene expression regulatory system responds in use in a transfected cell to said expression inducing agent which is provided by at least one of the following:

electromagnetic radiation, application of heat or cooling,
application of a magnetic or electric field, an exogenous
10 chemical inducing agent, radiation in the form of sub-atomic
particles.

26. Vector material as claimed in Claim 25 wherein said expression inducing agent is electromagnetic radiation in the form of ultra-violet or visible light.

15 27. Vector material as claimed in Claim 25 wherein said expression inducing influence is electromagnetic radiation in the form of X-rays or gamma-rays.

28. Vector material as claimed in Claim 25 wherein said expression inducing agent is electromagnetic radiation in the form of X-rays or gamma-
20 rays at a substantially sub-lethal dosage.

29. Vector material as claimed in Claim 23 or 24 wherein said expression inducing agent is an exogenous chemical expression inducing agent that induces cellular damage.

30. Vector material as claimed in Claim 23 or 24 wherein said expression
25 inducing agent is an exogenous chemical expression inducing agent in the form of an antitumour drug.

31. Vector material as claimed in Claim 30 wherein the antitumour drug is a platinum containing drug.

32. Vector material as claimed in Claim 30 wherein the antitumour drug is selected from the following:

5 Temozolomide, Dacarbazine, Streptozotocin, Procarbazine, Carmustine, Semustine, Lomustine, Fotemustine, Busulphan, Treosulphan, Mechlorethamine, Cyclophosphamide, Iphosphamide, Chlorambucil, Melphalan, ethyleneimines triethylene melamine, hexamethylmelamine, TEPA and thio-TEPA, dibromomannital and dibromodulcitol,

10 hydroxyurea, Methotrexate, azaserine Azathioprin, 5-azacytidine, 5-fluorouracil, cytosine arabinoside, 6-mercaptopurine, Allopurinol 6-thioguanine, deoxycytosine, Tiazofurin, Acivicin, Pyrazofurin and p-aminolaevulinic acid, plant alkaloids such as Vinblastine, Vincristine and Vindesine, Etoposide and Teniposide, antitumour antibiotics such as Doxorubicin, Daunorubicin, Actinomycin, Bleomycins, Mytomycin, Mythramycin, Mitozantrone hormones such as estrogen and progesterone.

15 33. Vector material as claimed in Claim 23 or 24 wherein said expression inducing agent is an exogenous chemical inducing agent in the form of a hormone that interacts with a receptor molecule which interacts with a complementary hormone responsive element in the control gene expression regulatory system.

20 34. Vector material as claimed in Claim 23 or 24 wherein the control gene expression regulatory system comprises a gene upregulation system that can be activated by a chemical agent.

25 35. Vector material as claimed in Claim 22 or 23 containing number of different control gene expression regulatory elements responsive to different

expression inducing influences so as to be activated under a range of different conditions.

36. Vector material as claimed in any of the preceding claims wherein at least one element of the sensitizing gene expression regulatory system is 5 inducible in response to the effect of a predetermined exogenous or endogenous expression inducing influence.

37. Vector material as claimed in Claim 36 wherein the sensitizing gene expression regulatory system includes an expression control element responsive in use in a transfected cell to a hypoxic condition in the cellular environment.

10 38. Vector material as claimed in any of the preceding claims wherein at least one element of the sensitizing gene expression regulatory system is selected for efficiency in the particular tumour(s) to which said antitumour therapy is directed, the selection being carried out using gene array technology.

15 39. Vector material as claimed in any of the preceding claims which includes a plurality of tumour sensitizing genes providing a range of different expression products.

20 40. Vector material as claimed in Claim 39 wherein the different expression products are prodrug activating agents which are effective in relation to a range of different prodrugs, or which provide a range of different immune system stimulatory factors.

41. A pharmaceutical composition comprising vector material as claimed in any of the preceding claims in association with a pharmaceutically acceptable carrier or excipient.

25 42. A pharmaceutical composition as claimed in Claim 41 which further includes a transfection agent.

43. A kit comprising one or more unit doses of vector material as defined in any one of Claims 1 to 40 together with a transfection agent.

44. A kit comprising:

- (a) a vector which comprises a tumour cell sensitizing gene or genes and a sensitizing gene expression regulatory system as defined in any one of the Claims 1 to 40;
- 5 (b) a vector which comprises a control gene and a control gene expression regulatory system as defined in any one of Claims 1 to 40; and
- (c) instructions for the use of vectors (a) and (b) in antitumour therapy.

45. A kit comprising:

- 10 (a) vector material as defined in any one of Claims 1 to 40;
- (b) a vector which comprises a tumour cell sensitizing gene or genes and a sensitizing gene expression regulatory system as defined in any one of Claims 1 to 40.

46. A kit as claimed in Claim 44 or 45 for use in antitumour therapy.

- 15 47. A kit as claimed in Claim 43, 44, 45 or 46 wherein each of the vectors and/or vector material is provided in the form of a pharmaceutical composition in association with a pharmaceutically acceptable carrier or excipient.

- 20 48. A kit as claimed in any one of Claims 43 to 47 wherein the sensitizing gene(s) produce(s) a prodrug activating agent and wherein one or more doses of a prodrug matched to said prodrug activating agent are also included in said kit.

49. A method of treatment for cancer patients wherein there is delivered to tumour cells vector material as claimed in any of Claims 1 to 40, said cells then being subjected to the appropriate expression inducing influence.

- 25 50. A method of treating tumour cells in a biological host system comprising:

(a) administering to the system an effective amount of a composition comprising vector material containing a tumour cell sensitizing gene or genes and having a control gene expression regulatory system responsive to a predetermined exogenous or endogenous expression inducing influence, said control gene expression regulatory system being operatively linked to a recombinase gene, together with recombinase target sites flanking a region of which removal permits continuous expression of said tumour cell sensitizing gene(s) as specified above, or primes said sensitizing gene(s) for continuous induced expression;

(b) causing said tumour cells transfected with said vector material to be subjected to a dose of an expression inducing agent effective to activate the recombinase gene expression control system of said vector material, thereby to bring about, via recombinase-mediated site specific recombination within the vector material, expression of the or each tumour cell sensitizing gene component;

and, in the case of prodrug activating sensitizing genes,

(c) administering to the host an effective amount of a composition comprising a prodrug convertible into an active form by the expression product of said tumour cell sensitizing gene or genes.

51. Use of vector material as defined in any one of Claims 1 to 40 in the manufacture of a medicament or of a kit as defined in Claim 43 or 45 for use in antitumour therapy.

52. Use of a vector which comprises a tumour cell sensitizing gene or genes and a sensitizing gene expression regulatory system as defined in any one of Claims 1 to 40 in the manufacture of a kit as defined in Claim 44 for use in antitumour therapy.

53. Use of a vector which comprises a control gene and a control gene expression regulatory system as defined in any one of Claims 1 to 40 in the manufacture of a kit as defined in Claim 44 or 45 for use in antitumour therapy.

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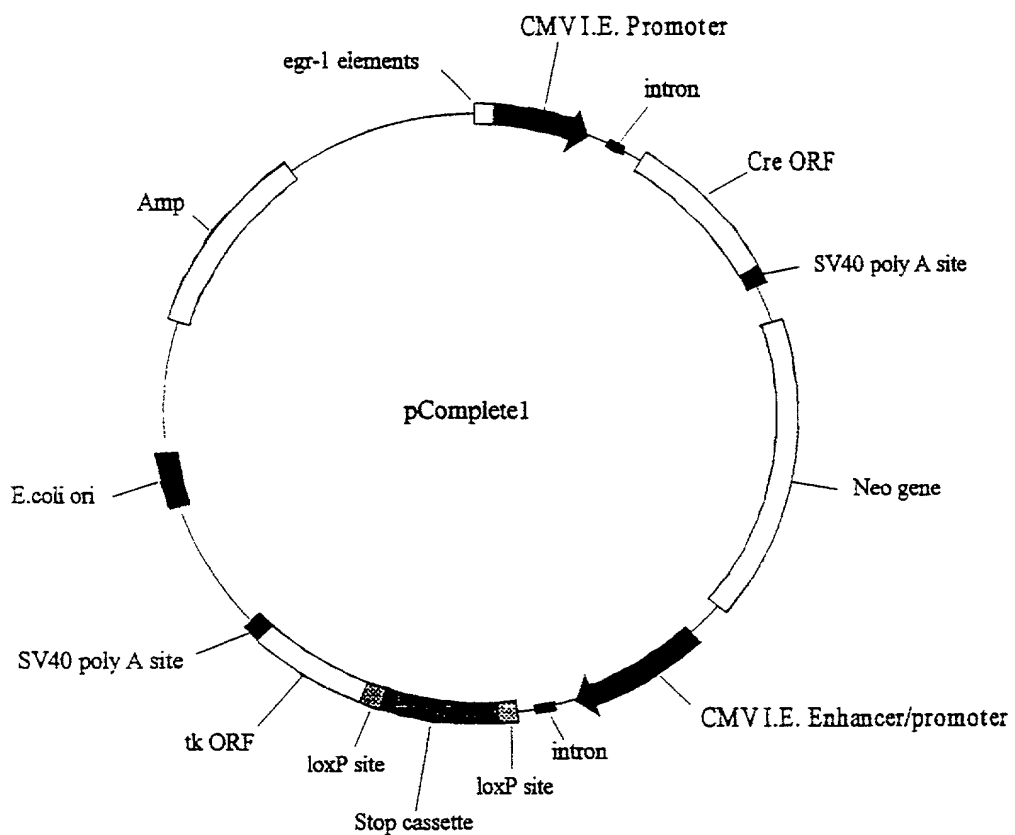


Figure 1.

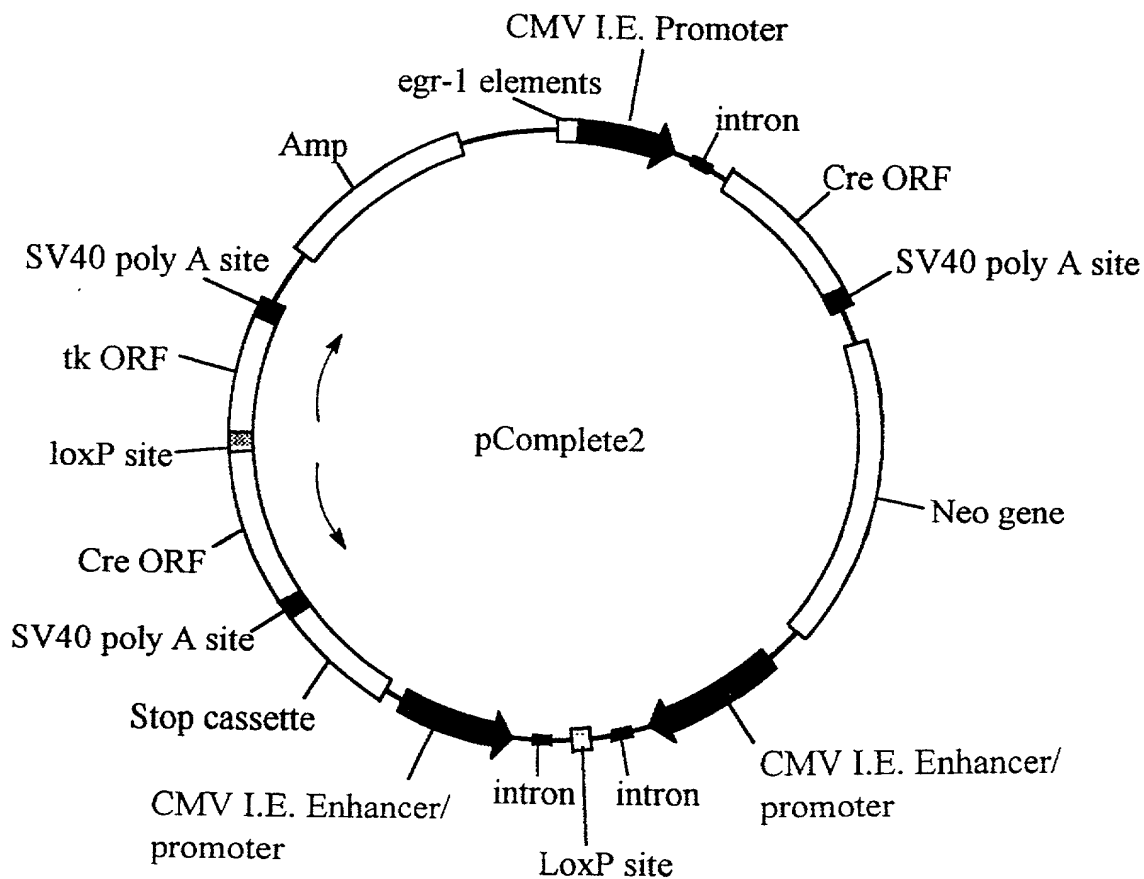


Figure 2.

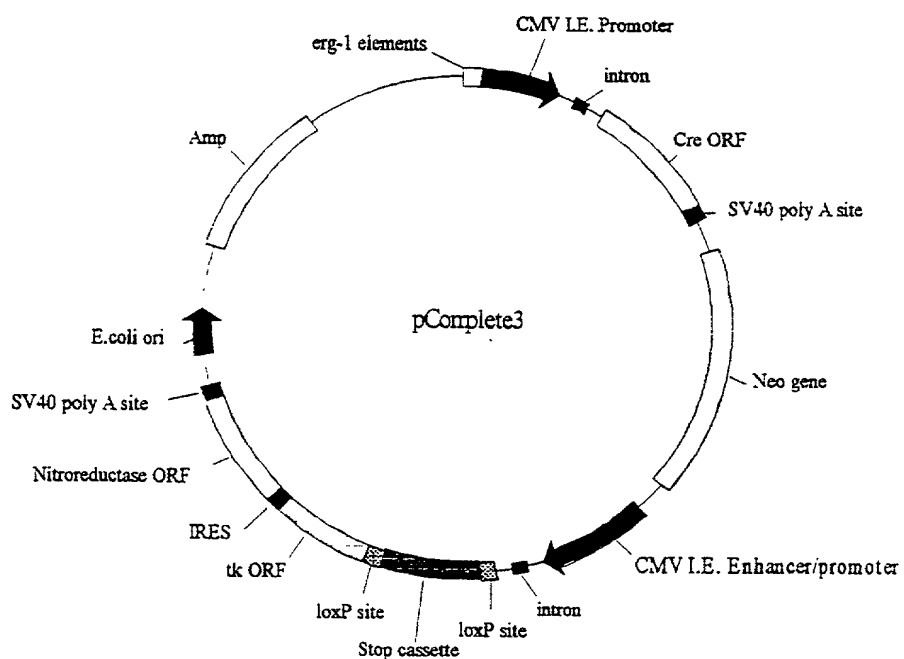


Figure 3.

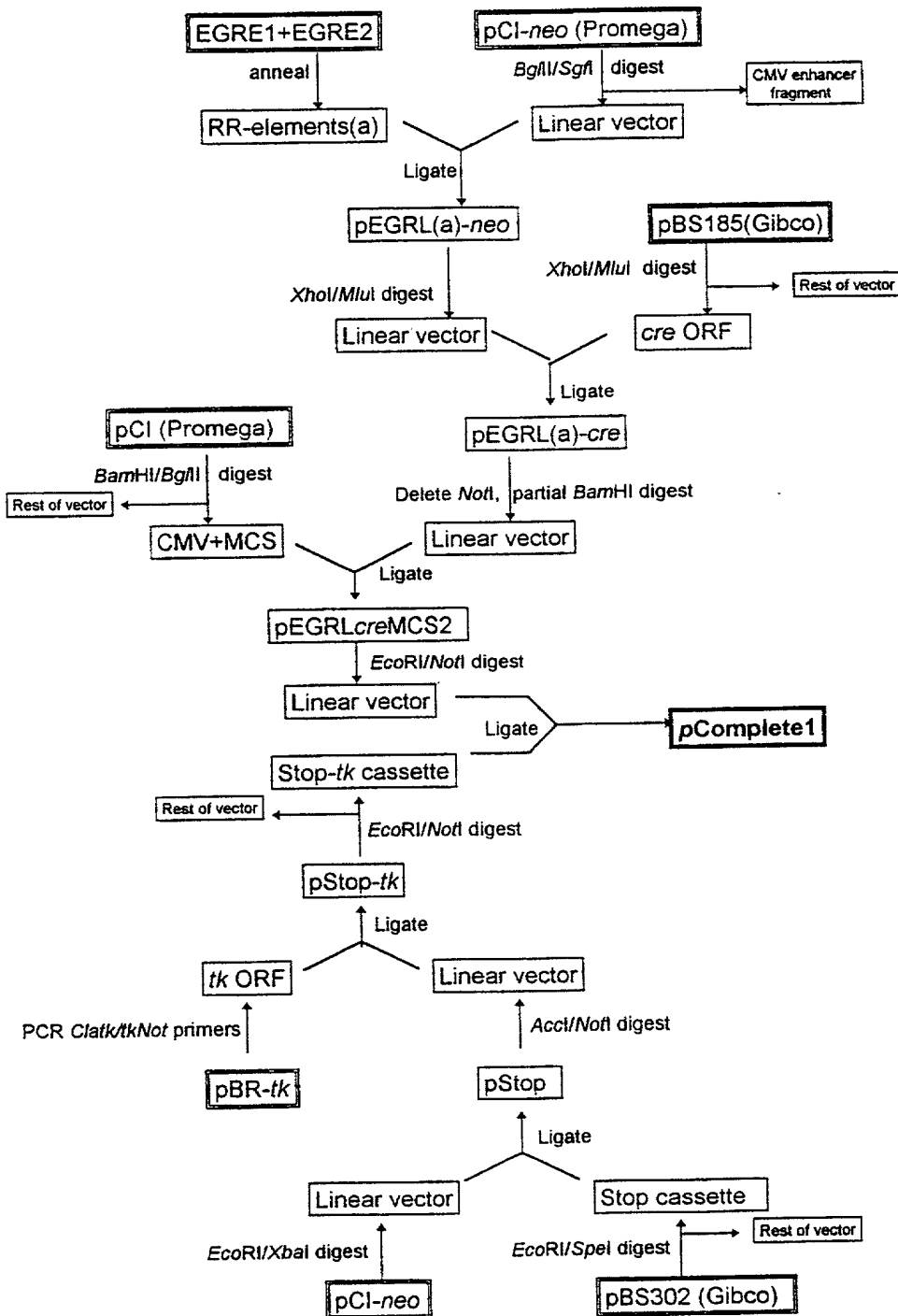


Figure 4.

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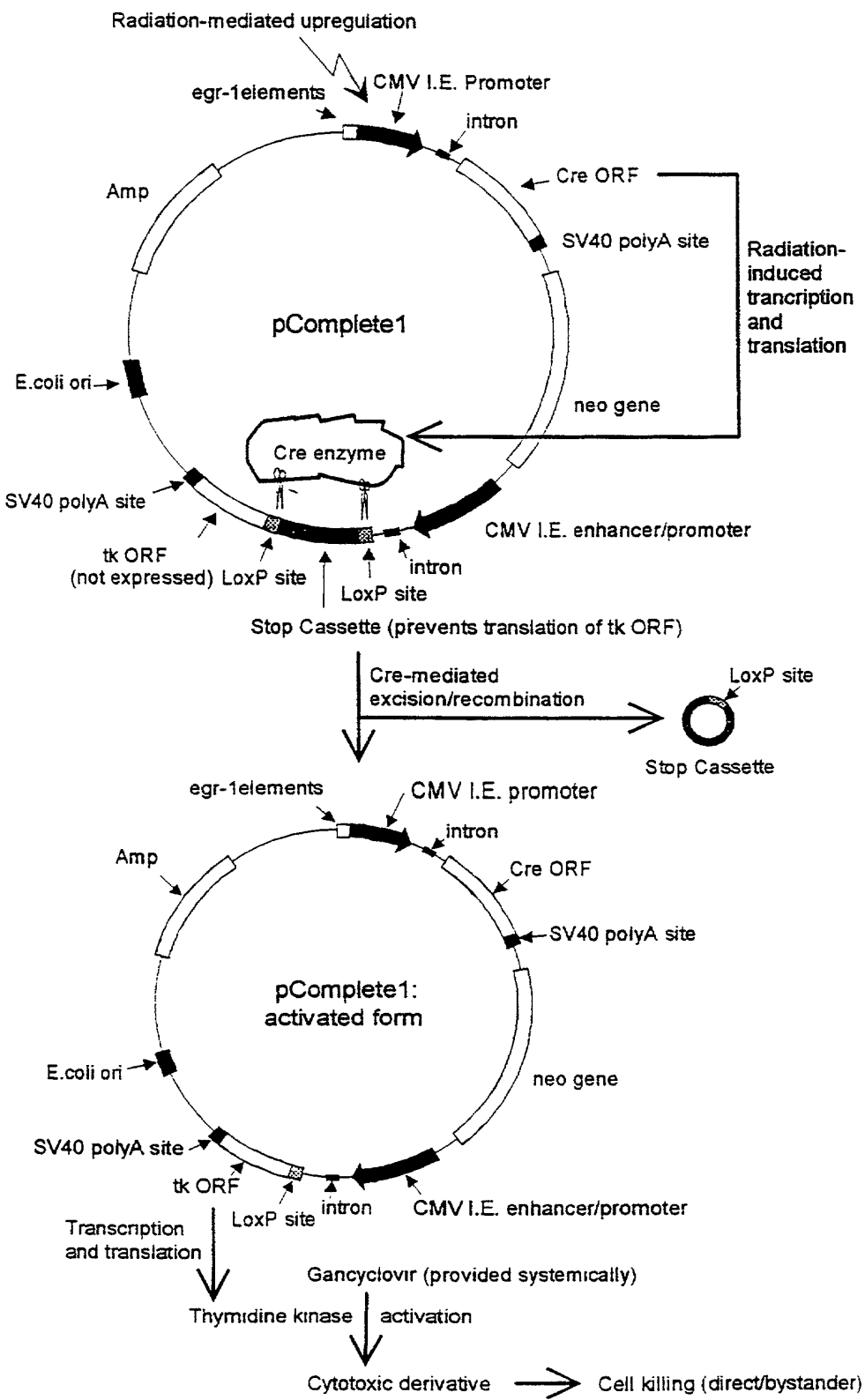


Figure 5.

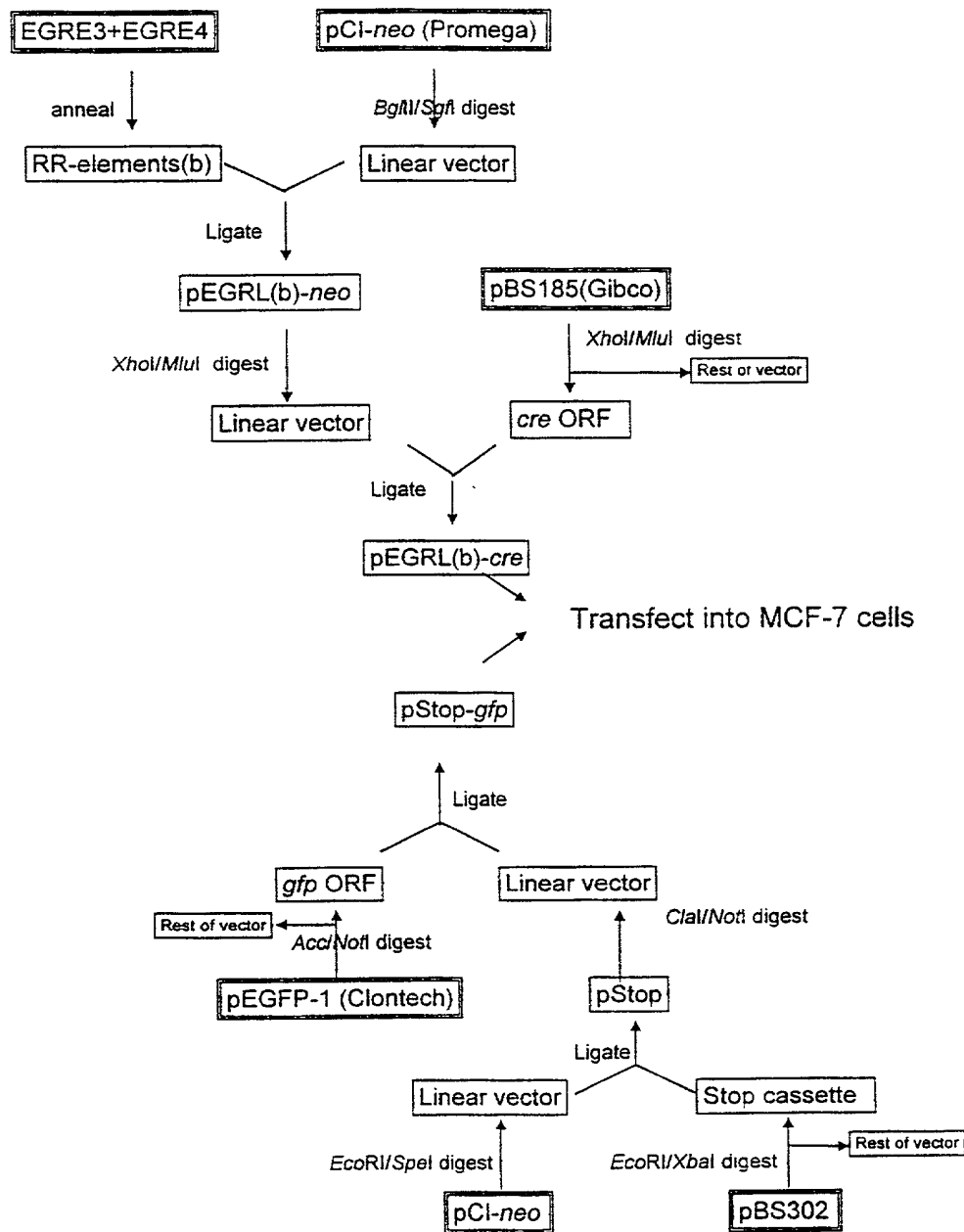


Figure 6.

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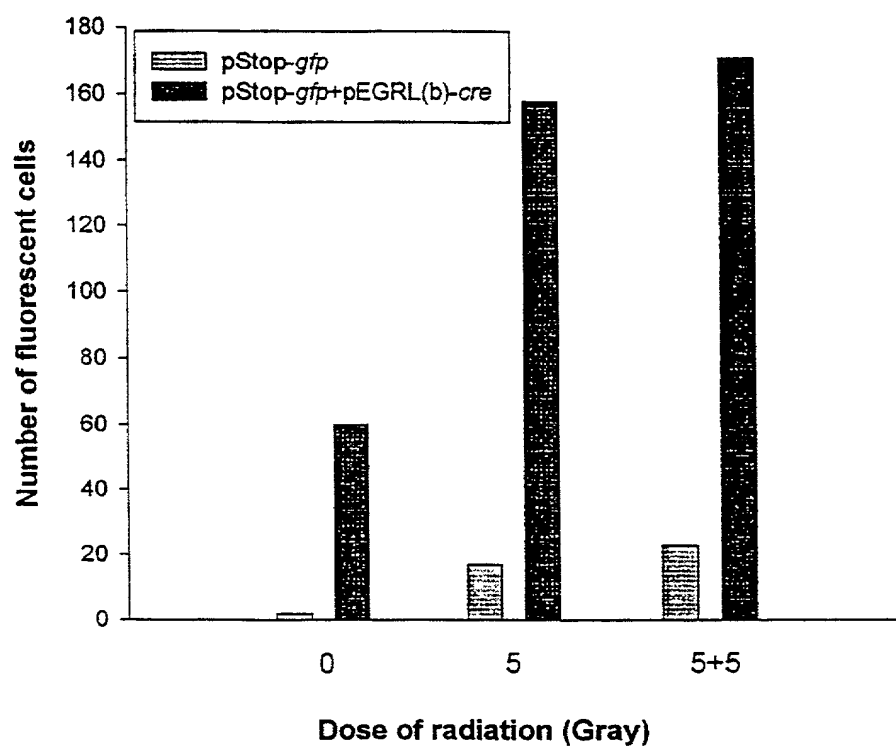


Figure 7.

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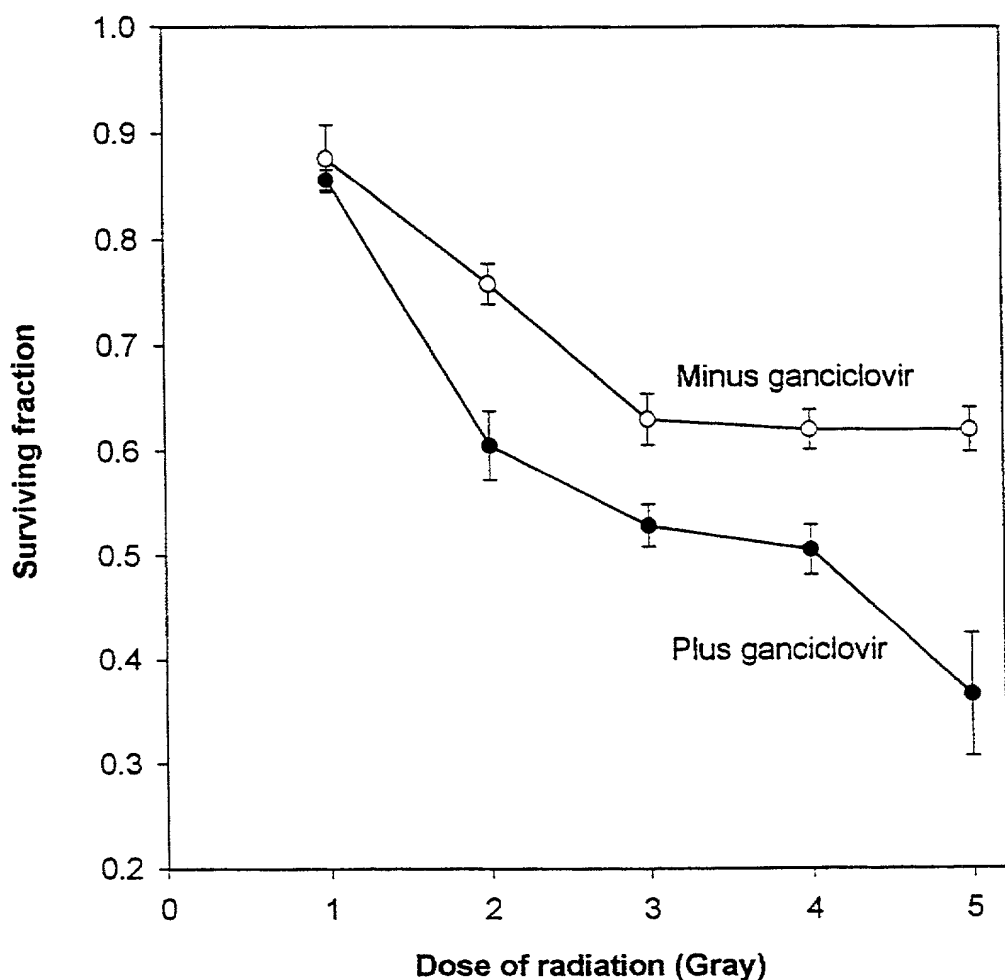


Figure 8.

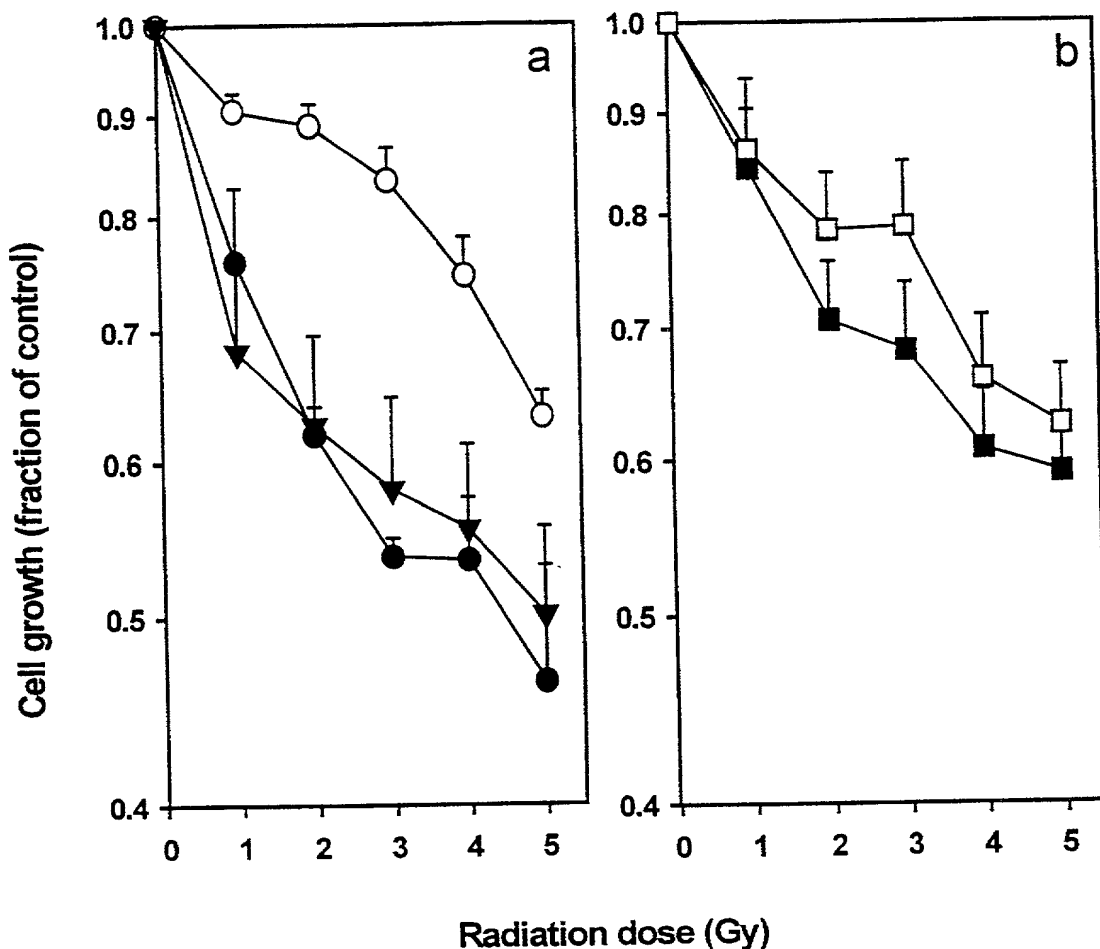


Figure 9a and 9b

10/11

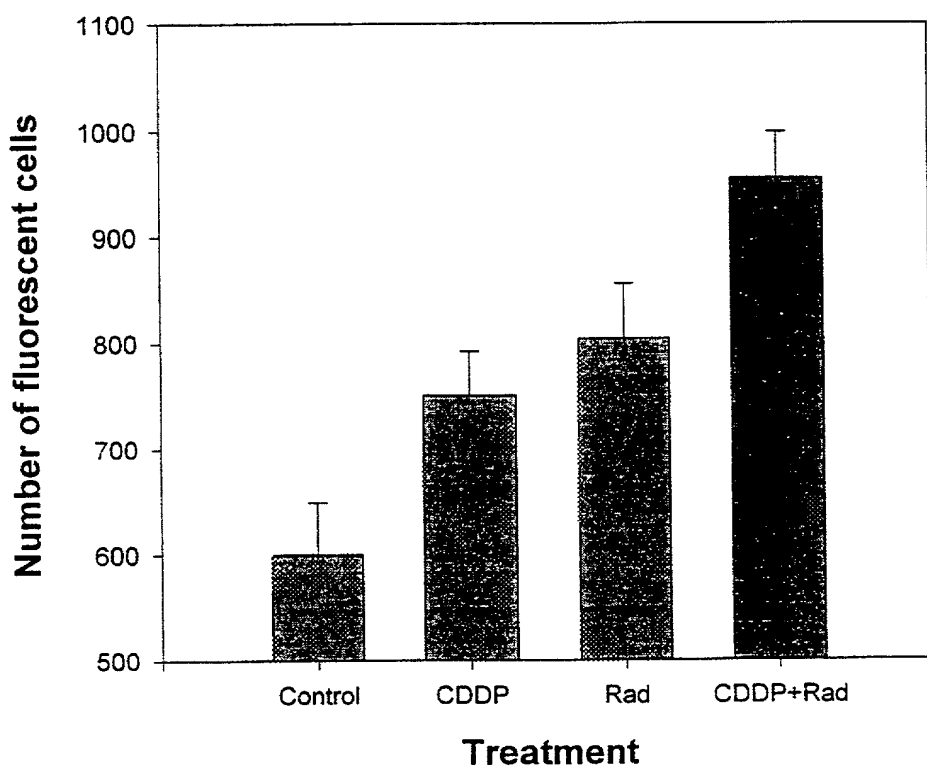


Figure 10.

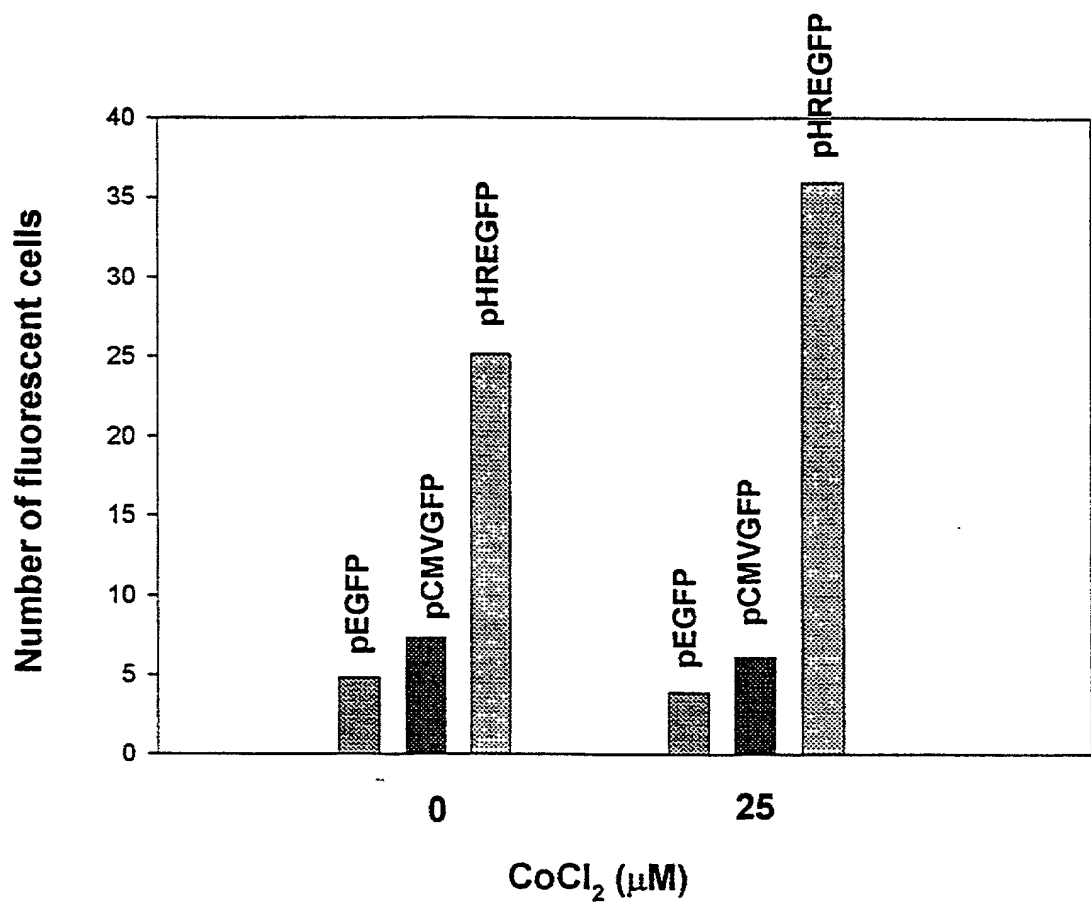


Figure 11.

FOR UTILITY/DESIGN
CIP/PCT NATIONAL/PLANT
ORIGINAL/SUBSTITUTE/SUPPLEMENTAL
DECLARATIONS

RULE 63 (37 C.F.R. 1.63)
DECLARATION AND POWER OF ATTORNEY
FOR PATENT APPLICATION
IN THE UNITED STATES PATENT AND TRADEMARK OFFICE

PM&S
FORM

I, a below named inventor, I hereby declare that my residence, post office address and citizenship are as stated below next to my name, and I believe I am the original, first and sole inventor (if only one name is listed below) or an original, first and joint inventor (if plural names are listed below) of the subject matter which is claimed and for which a patent is sought on the INVENTION ENTITLED
GENE THERAPY VECTORS AND THEIR USE IN ANTITUMOUR THERAPY

the specification of which (CHECK applicable BOX(ES))

X → [] is attached hereto.
BOX(ES) → [X] was filed on November 14, 2000 as U.S. Application No. 09/700,259
→ [] was filed as PCT International Application No. PCT/GB99/01362 on 17 MAY 1999
→ & (if U.S. or PCT application amended) was amended on

I hereby state that I have reviewed and understand the contents of the above identified specification, including the claims, as amended by any amendment referred to above. I acknowledge the duty to disclose all information known to me to be material to patentability as defined in 37 C.F.R. 1.56. I hereby claim foreign priority benefits under 35 U.S.C. 119/365 of any foreign application(s) for patent or inventor's certificate listed below and have also identified below any foreign application for patent or inventor's certificate filed by me or my assignee disclosing the subject matter claimed in this application and having a filing date (1) before that of the application on which priority is claimed, or (2) if no priority claimed, before the filing date of this application:

PRIOR FOREIGN APPLICATION(S)		Date first Laid-open or Published	Date Patented or Granted	Priority Claimed
Number	Country			Yes No
9810423.5	UNITED KINGDOM	15 MAY 1998		X
PCT/GB99/01362	PCT	17 MAY 1999		X

I hereby claim domestic priority benefit under 35 U.S.C. 119/120/365 of the indicated United States applications listed below and PCT international applications listed above or below and, if this is a continuation-in-part (CIP) application, insofar as the subject matter disclosed and claimed in this application is in addition to that disclosed in such prior applications, I acknowledge the duty to disclose all information known to me to be material to patentability as defined in 37 C.F.R. 1.56 which became available between the filing date of each such prior application and the national or PCT international filing date of this application:

PRIOR U.S. PROVISIONAL, NONPROVISIONAL AND/OR PCT APPLICATION(S)		Status	Priority Claimed
Application No. (series code/serial no.)	Dav/Month/Year Filed		Yes No
PCT/GB99/01362	17 MAY 1999	PENDING	X

I hereby declare that all statements made herein of my own knowledge are true and that all statements made on information and belief are believed to be true; and further that these statements were made with the knowledge that willful false statements and the like so made are punishable by fine or imprisonment, or both, under Section 1001 of Title 18 of the United States Code and that such willful false statements may jeopardize the validity of the application or any patent issued thereon.

And I hereby appoint Pillsbury Madison & Sutro LLP, Intellectual Property Group, 1100 New York Avenue, N.W., Ninth Floor, East Tower, Washington, D.C. 20005-3918, telephone number (202) 861-3000 (to whom all communications are to be directed), and the below-named persons (of the same address) individually and collectively my attorneys to prosecute this application and to transact all business in the Patent and Trademark Office connected therewith and with the resulting patent, and I hereby authorize them to delete names/numbers below of persons no longer with their firm and to act and rely on instructions from and communicate directly with the person/assignee/attorney/firm/ organization who/which first sends/sent this case to them and by whom/which I hereby declare that I have consented after full disclosure to be represented unless/until I instruct the above Firm and/or a below attorney in writing to the contrary.

Paul N. Kokulis	16723	George M. Sirilla	18221	G. Paul Edgell	24238	Richard H. Zaitlen	27248
Raymond F. Lippitt	17519	Donald J. Bird	25323	Lynn E. Eccleston	35861	Roger R. Wise	31204
G. Lloyd Knight	17698	Peter W. Gowdey	25872	David A. Jakopin	32995	Jay M. Finkelstein	21082
Carl G. Love	18781	Dale S. Lazar	28872	Mark G. Paulson	30793	Anita M. Kirkpatrick	32617
Edgar H. Martin	20534	Glenn J. Perry	28458	Timothy J. Klima	34852	Michael R. Dzwonczyk	36787
William K. West, Jr.	22057	Kendrew H. Colton	30368	Stephen C. Glazier	31361		
Kevin E. Joyce	20508	Paul E. White, Jr.	32011	Paul F. McQuade	31542		
David W. Brinkman	20817	Michelle N. Lester	32331	Ruth N. Morduch	31044		

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(FOR ADDITIONAL INVENTORS, check box [X] and attach sheet (PAT-116.2) for same information for each re signature, name, date, citizenship, residence and address.)

DECLARATION AND POWER OF ATTORNEY

(continued)

ADDITIONAL INVENTORS.

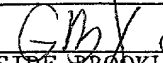
Page 2

INVENTOR'S SIGNATURE: Date 14/6/01Inventor's Name (typed) JOLYONFirst Initial H Middle Initial G Family Name HENDRY

UNITED KINGDOM

Residence (City) CHESHIRE(State/Foreign Country) UNITED KINGDOM

Country of Citizenship

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Date

Inventor's Name (typed)

First

Middle Initial

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(State/Foreign Country)

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6. INVENTOR'S SIGNATURE: 

Date

Inventor's Name (typed)

First

Middle Initial

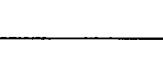
Family Name

Country of Citizenship

Residence (City)

(State/Foreign Country)

Post Office Address (Include Zip Code)

7. INVENTOR'S SIGNATURE: 

Date

Inventor's Name (typed)

First

Middle Initial

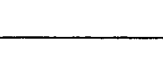
Family Name

Country of Citizenship

Residence (City)

(State/Foreign Country)

Post Office Address (Include Zip Code)

8. INVENTOR'S SIGNATURE: 

Date

Inventor's Name (typed)

First

Middle Initial

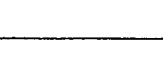
Family Name

Country of Citizenship

Residence (City)

(State/Foreign Country)

Post Office Address (Include Zip Code)

9. INVENTOR'S SIGNATURE: 

Date

Inventor's Name (typed)

First

Middle Initial

Family Name

Country of Citizenship

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(State/Foreign Country)

Post Office Address (Include Zip Code)

10. INVENTOR'S SIGNATURE: 

Date

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11. INVENTOR'S SIGNATURE: 

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Middle Initial

Family Name

Country of Citizenship

Residence (City)

(State/Foreign Country)

Post Office Address (Include Zip Code)

FOR ADDITIONAL INVENTORS, check box and attach sheet with same information and signature and date for each.

SEQUENCE LISTING

5 (1) GENERAL INFORMATION:

(i) APPLICANT:

- (A) NAME: CANCER RESEARCH CAMPAIGN TECHNOLOGY LIMITED
- (B) STREET: CAMBRIDGE HOUSE, 6-10 CAMBRIDGE TERRACE,
REGENT'S PARK,
- (C) CITY: LONDON
- (E) COUNTRY: UNITED KINGDOM
- (F) POSTAL CODE (ZIP): NW1 4JL

15 (ii) TITLE OF INVENTION:
IONIZING RADIATION OR DIATHERMY-SWITCHED GENE
THERAPY VECTORS AND THEIR USE IN ANTITUMOUR THERAPY

(iii) NUMBER OF SEQUENCES: 12

(iv) COMPUTER READABLE FORM:

- (A) MEDIUM TYPE: Floppy disk
- (B) COMPUTER: IBM PC compatible
- (C) OPERATING SYSTEM: PC-DOS/MS-DOS
- (D) SOFTWARE: PatentIn Release #1.0, Version #1.30 (EPO)

30 (2) INFORMATION FOR SEQ ID NO: 1:

(i) SEQUENCE CHARACTERISTICS:

- (A) LENGTH: 10 base pairs
- (B) TYPE: nucleic acid
- (C) STRANDEDNESS: single
- (D) TOPOLOGY: linear

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 1:

40 CCTTATTTGG

10

45 (2) INFORMATION FOR SEO ID NO: 2:

(i) SEQUENCE CHARACTERISTICS:

- (A) LENGTH: 69 base pairs
- (B) TYPE: nucleic acid
- (C) STRANDEDNESS: single
- (D) TOPOLOGY: linear

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 2:

55 GATCTCCTTA TTTGGCCTTA TTTGGCCTTA TTTGGCCTTA TTTGGCCTTA TTTGGCCTTA TTTGGCCTTA 60
TTGGGCGAT 69

60

(2) INFORMATION FOR SEQ ID NO: 3:

(i) SEQUENCE CHARACTERISTICS:

5 (A) LENGTH: 64 base pairs
(B) TYPE: nucleic acid
(C) STRANDEDNESS: single
(D) TOPOLOGY: linear

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 3:

10 CGCCCAAATA AGGCCAAATA AGGCCAAATA AGGCCAAATA AGGCCAAATA AGGCCAAATA 60
AGGA 64

15

(2) INFORMATION FOR SEQ ID NO: 4:

(i) SEQUENCE CHARACTERISTICS:

20 (A) LENGTH: 48 base pairs
(B) TYPE: nucleic acid
(C) STRANDEDNESS: single
(D) TOPOLOGY: linear

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 4:

25 GATCTTTATT TGGCCTTATT TGGCCTTATT TGGCCTTATT TGGGCGAT 48

30

(2) INFORMATION FOR SEQ ID NO: 5:

(i) SEQUENCE CHARACTERISTICS:

35 (A) LENGTH: 44 base pairs
(B) TYPE: nucleic acid
(C) STRANDEDNESS: single
(D) TOPOLOGY: linear

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 5:

40 CGCCCAAATA AGGCCAAATA AGGCCAAATA AGGCCAAATA AGGA 44

45

(2) INFORMATION FOR SEQ ID NO: 6:

(i) SEQUENCE CHARACTERISTICS:

50 (A) LENGTH: 36 base pairs
(B) TYPE: nucleic acid
(C) STRANDEDNESS: single
(D) TOPOLOGY: linear

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 6:

55 TCCAGATCTC CCGGTTCGCT CTCACGGTCC CTGAGG 36

60

(2) INFORMATION FOR SEQ ID NO: 7:

(i) SEQUENCE CHARACTERISTICS:

- (A) LENGTH: 32 base pairs
- (B) TYPE: nucleic acid
- (C) STRANDEDNESS: single
- (D) TOPOLOGY: linear

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 7:

10 CGGCGCGCCG CTGGATCTCT CGCGACTCCCC CG

32

15

(2) INFORMATION FOR SEQ ID NO: 8:

(i) SEQUENCE CHARACTERISTICS:

- (A) LENGTH: 42 base pairs
- (B) TYPE: nucleic acid
- (C) STRANDEDNESS: single
- (D) TOPOLOGY: linear

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 8:

25 ACTGCGATCG CGGGCCCCGGC CCGGCCCGCA TCCCAGGGCCC CC

42

30

(2) INFORMATION FOR SEQ ID NO: 9:

(i) SEQUENCE CHARACTERISTICS:

- (A) LENGTH: 26 base pairs
- (B) TYPE: nucleic acid
- (C) STRANDEDNESS: single
- (D) TOPOLOGY: linear

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 9:

40 CCATCGATAT GGCTTCGTAC CCCGGC

26

45

(2) INFORMATION FOR SEQ ID NO: 10:

(i) SEQUENCE CHARACTERISTICS:

50 (A) LENGTH: 40 base pairs
(B) TYPE: nucleic acid
(C) STRANDEDNESS: single
(D) TOPOLOGY: linear

55 (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 10:

AAGGAAAAAA GCGGCCGCCT CCTTCCGTGT TTCAGTTAGC

40

(2) INFORMATION FOR SEQ ID NO: 11:

(i) SEQUENCE CHARACTERISTICS:

5 (A) LENGTH: 83 base pairs
 (B) TYPE: nucleic acid
 (C) STRANDEDNESS: single
 (D) TOPOLOGY: linear

10 (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 11:

GATCTAGGGC CGGACGTGGG GCCCCGTAGG CACGCTGAGT GCGTGCGGGA 50

CTCGGAGTAC GTGACGGAGC CCCGCGATGC GAT 83

15

20 (2) INFORMATION FOR SEQ ID NO: 12:

(i) SEQUENCE CHARACTERISTICS:

25 (A) LENGTH: 77 base pairs
 (B) TYPE: nucleic acid
 (C) STRANDEDNESS: single
 (D) TOPOLOGY: linear

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 12:

CGCATCGCGG GGCTCCGTCA CGTACTCCGA GTCCCCGCACG CACTCAGCGT GCCTACGGGG 60

CCCCACGTCC GGCCCTTA